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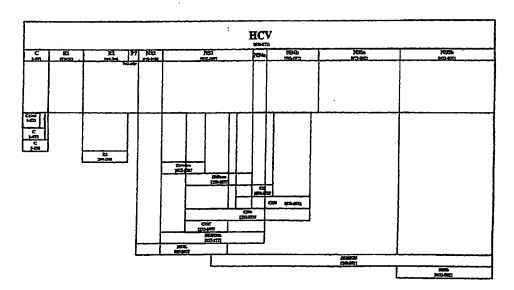
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[Continued on next page]

(54) Title: ACTIVATION OF HCV-SPECIFIC CELLS

HCV Genome and Recombinant Proteins



(57) Abstract: The invention provides a method of activating hepatitis C virus (HCV)-specific T cells, including CD4* and CD8* T cells. HCV-specific T cells are activated using fusion proteins comprising HCV NS3, NS4, NS5a, and NS5b polypeptides, polynucleotides encoding such fusion proteins, or polypeptide or polynucleotide compositions containing the individual components of these fusions. The method can be used in model systems to develop HCV-specific immunogenic compositions, as well as to immunize a mammal against HCV.

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ACTIVATION OF HCV-SPECIFIC T CELLS

TECHNICAL AREA OF THE INVENTION

The invention relates to the activation of hepatitis C virus(HCV)-specific T cells. More particularly, the invention relates to the use of multiple HCV polypeptides, either alone or as fusions, to stimulate cell-mediated immune responses, such as to activate HCV-specific T cells.

15 BACKGROUND OF THE INVENTION

Hepatitis C virus (HCV) infection is an important health problem with approximately 1% of the world's population infected with the virus. Over 75% of acutely infected individuals eventually progress to a chronic carrier state that can result in cirrhosis, liver failure, and hepatocellular carcinoma. See Alter et al. (1992) N. Engl. J. Med. 327:1899-1905; Resnick and Koff. (1993) Arch. Intem. Med. 153:1672-1677; Seeff (1995) Gastrointest. Dis. 6:20-27; Tong et al. (1995) N. Engl. J. Med. 332:1463-1466.

Despite extensive advances in the development of pharmaceuticals against certain viruses like HIV, control of acute and chronic HCV infection has had limited success (Hoofnagle and di Bisceglie (1997) N. Engl. J. Med. 336:347-356). In particular, generation of a strong cytotoxic T lymphocyte (CTL) response is thought to be important for the control and eradication of HCV infections. Thus, there is a need in the art for effective methods of inducing strong CTL responses against HCV.

SUMMARY OF THE INVENTION

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It is an object of the invention to provide reagents and methods for stimulating immune responses, such as activating T cells which recognize epitopes of HCV polypeptides. This and other objects of the invention are provided by one or more of the embodiments described below.

The invention provides HCV proteins useful for stimulating immune responses, such as activating HCV-specific T cells. One embodiment provides a fusion protein that comprises HCV polypeptides, wherein the HCV polypeptides consist essentially of an NS3, an NS4, an NS5a polypeptide, and optionally a core polypeptide. In certain embodiments, the fusion protein includes an NS5b polypeptide.

In certain embodiments, at least one of the HCV polypeptides is derived from a different strain of HCV than the other polypeptides.

The invention also provides compositions comprising any of these fusion proteins and a pharmaceutically acceptable excipient. In certain embodiments, the compositions further comprise an adjuvant, a CpG polynucleotide and/or the fusion protein is adsorbed to or entrapped within a microparticle or ISCOM. The compositions can further comprise a polynucleotide encoding an E1E2 complex. The E1E2 polynucleotide can also be adsorbed to or entrapped withing a microparticle.

Another embodiment provides a composition comprising HCV polypeptides and a pharmaceutically acceptable excipient. The HCV polypeptides consist essentially of an NS3, an NS4, an NS5a polypeptide, and optionally a core polypeptide. In certain embodiments, the composition includes an NS5b polypeptide. In other embodiments, the compositions further comprise an adjuvant, a CpG polynucleotide and/or one or more of the HCV polypeptides is adsorbed to or entrapped within a microparticle or ISCOM. The compositions can further comprise a polynucleotide encoding an E1E2 complex. The E1E2 polynucleotide can also be adsorbed to or entrapped withing a microparticle.

Moreover, one of the HCV polypeptides may be derived from a different strain of HCV than the others.

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Even another embodiment of the invention provides an isolated and purified polynucleotide which encodes a fusion protein as described above. In additional embodiments, the fusion proteins further include a polynucleotide encoding an E1E2 complex.

Yet another embodiment of the invention provides a composition comprising the polynucleotides described above and a pharmaceutically acceptable excipient. In certain embodiments, the compositions further comprise an adjuvant and/or the polynucleotide may be adsorbed to or entrapped within a microparticle. The compositions can further comprise a polynucleotide encoding an E1E2 complex. The E1E2 polynucleotide can also be adsorbed to or entrapped withing a microparticle.

In a further embodiment, the invention provides a composition comprising HCV polynucleotides and a pharmaceutically acceptable excipient, wherein the HCV polynucleotides consist essentially of polynucleotides encoding an NS3, an NS4, an NS5a polypeptide, and optionally a core polypeptide. In certain embodiments, the composition also includes a polynucleotide encoding an NS5b polypeptide. The compositions may further comprise an adjuvant and/or one or more of the polynucleotides may be adsorbed to or entrapped within a microparticle. The compositions can further comprise a polynucleotide encoding an E1E2 complex. The E1E2 polynucleotide can also be adsorbed to or entrapped withing a microparticle. Additionally, one or more of the polynucleotides may be derived from a different strain of HCV than the others.

In another embodiment, the invention provides a method of activating T cells which recognize an epitope of an HCV polypeptide. T cells are contacted with any of the fusions, polynucleotides or compositions described above. A population of activated T cells recognizes an epitope of the NS3, NS4, NS5a, NS5b, core and/or E1E2 polypeptide.

In the proteins and polynucleotides above, the regions in the fusions need not be in the order in which they naturally occur in the native HCV polyprotein. Thus, for example, the NS5b polypeptide, if present, may be at the N- and/or C-terminus of the fusion, or may be located internally. Similarly, the E1 polypeptide may precede or follow the E2 polypeptide. The E1E2 polypeptide may also be part of the

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nonstructural fusion protein or may be provided separately, as an E1E2 complex, or as individual polypeptides.

Moreover, the NS3 polypeptide may include a modification to inhibit protease activity, such that cleavage of the fusion is inhibited. Such modifications are described more fully below. Additionally, the compositions can comprise more than one HCV nonstructural fusion protein, such as a fusion protein with NS3, NS4 and NS5a, and a fusion protein with NS3, NS4, NS5a, NS5b and E1E2. The E1E2 complexes, whether present separately or as part of the fusion, can have varying E1E2 polypeptides (described more fully below).

In certain embodiments, the nonstructural fusion protein consists of, from the amino terminus to the carboxyl terminus, an NS3, an NS4, an NS5a and, optionally, an NS5b polypeptide and the E1E2 complex consists of, from amino terminus to the carboxyl terminus, an E1 polypeptide and an E2 polypeptide.

The various polypeptides (and polynucleotides encoding therefor) are derived from the same HCV isolate, or from different strains and isolates including isolates having any of the various HCV genotypes, to provide increased protection against a broad range of HCV genotypes.

Yet another embodiment of the invention provides a method of stimulating an immune response, such as a cellular immune response, in a vertebrate subject by administering a composition as described herein. In certain embodiments, the composition activates T cells which recognize an epitope of an HCV polypeptide. T cells are contacted with a composition as described above. A population of activated T cells recognizes an epitope of one or more of the HCV polypeptide(s).

The invention thus provides methods and reagents for stimulating immune responses to HCV, such as for activating T cells which recognize epitopes of HCV polypeptides. These methods and reagents are particularly advantageous for identifying epitopes of HCV polypeptides associated with a strong CTL response and for immunizing mammals, including humans, against HCV.

BRIEF DESCRIPTION OF THE FIGURES

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Figure 1 is a diagrammatic representation of the HCV genome, depicting the various regions of the HCV polyprotein.

Figure 2 (SEQ ID NOS: 9 and 10) depicts the DNA and corresponding amino acid sequence of a representative native NS3 protease domain.

Figures 3A-3C (SEQ ID NOS:3 and 4) shows the nucleotide and corresponding amino acid sequence for the HCV-1 E1/E2/p7 region. The numbers shown in the figure are relative to the full-length HCV-1 polyprotein. The E1, E2 and p7 regions are shown.

Figure 4 is a diagram of plasmid pMHE1E2-809, encoding E1E2₈₀₉, a representative E1E2 protein for use with the present invention.

Figures 5A-5J (SEQ ID NOS:7 and 8) depict the DNA and corresponding amino acid sequence of a representative NS345Core fusion protein. The depicted sequence includes amino acids 1242-3011 of the HCV polyprotein (representing polypeptides from NS3, NS4, NS5a and NS5b) with amino acids 1-121 of the HCV polyprotein (representing a polypeptide from the core region) fused to the C-terminus of NS5b. This numbering is relative to the HCV-1 polyprotein.

Figure 6 shows a side-by-side comparison of IFN-γ expression generated in animals in response to delivery of alphavirus constructs encoding NS3NS4NS5a.

Figure 7 shows IFN-γ expression generated in animals in response to delivery of plasmid DNA encoding NS3NS4NS5a ("naked"), PLG-linked DNA encoding NS3NS4NS5a ("PLG), separate DNA plasmids encoding NS5a, NS34a, and NS4ab ("naked"), and PLG-linked DNA encoding NS5a, NS34a, and NS4ab ("PLG").

Figure 8 shows HCV-specific CD8+ and CD4+ responses in vaccinated chimpanzees.

Figure 9 depicts the specificity of T cell responses primed by electroporation of plasmid DNA two weeks subsequent to the third immunization.

Figure 10 shows the specificity of T cell responses primed by vaccinating chimpanzees with NS345Core₁₂₁-ISCOMS two weeks subsequent to the third immunization.

DETAILED DESCRIPTION OF THE INVENTION

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The practice of the present invention will employ, unless otherwise indicated, conventional methods of chemistry, biochemistry, recombinant DNA techniques and immunology, within the skill of the art. Such techniques are explained fully in the literature. See, e.g., Sambrook, et al., *Molecular Cloning: A Laboratory Manual* (2nd Edition); *Methods In Enzymology* (S. Colowick and N. Kaplan eds., Academic Press, Inc.); *DNA Cloning*, Vols. I and II (D.N. Glover ed.); *Oligonucleotide Synthesis* (M.J. Gait ed.); *Nucleic Acid Hybridization* (B.D. Hames & S.J. Higgins eds.); *Animal Cell Culture* (R.K. Freshney ed.); Perbal, B., *A Practical Guide to Molecular Cloning*.

It must be noted that, as used in this specification and the appended claims, the singular forms "a", "an" and "the" include plural referents unless the content clearly dictates otherwise. Thus, for example, reference to "an antigen" includes a mixture of two or more antigens, and the like.

The following amino acid abbreviations are used throughout the text:

15	Alanine: Ala (A)	Arginine: Arg (R)
	Asparagine: Asn (N)	Aspartic acid: Asp (D)
	Cysteine: Cys (C)	Glutamine: Gln (Q)
	Glutamic acid: Glu (E)	Glycine: Gly (G)
•	Histidine: His (H)	Isoleucine: Ile (I)
20	Leucine: Leu (L)	Lysine: Lys (K)
	Methionine: Met (M)	Phenylalanine: Phe (F)
	Proline: Pro (P)	Serine: Ser (S)
	Threonine: Thr (T)	Tryptophan: Trp (W)
	Tyrosine: Tyr (Y)	Valine: Val (V)

I. Definitions

In describing the present invention, the following terms will be employed, and are intended to be defined as indicated below.

The terms "polypeptide" and "protein" refer to a polymer of amino acid residues and are not limited to a minimum length of the product. Thus, peptides, oligopeptides, dimers, multimers, and the like, are included within the definition.

Both full-length proteins and fragments thereof are encompassed by the definition. The terms also include postexpression modifications of the polypeptide, for example, glycosylation, acetylation, phosphorylation and the like. Furthermore, for purposes of the present invention, a "polypeptide" refers to a protein which includes modifications, such as deletions, additions and substitutions (generally conservative in nature), to the native sequence, so long as the protein maintains the desired activity. These modifications may be deliberate, as through site-directed mutagenesis, or may be accidental, such as through mutations of hosts which produce the proteins or errors due to PCR amplification.

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10 An HCV polypeptide is a polypeptide, as defined above, derived from the HCV polyprotein. The polypeptide need not be physically derived from HCV, but may be synthetically or recombinantly produced. Moreover, the polypeptide may be derived from any of the various HCV strains and isolates including isolates having any of the 6 genotypes of HCV described in Simmonds et al., J. Gen. Virol. (1993) 15 74:2391-2399 (e.g., strains 1, 2, 3, 4 etc.), as well as newly identified isolates, and subtypes of these isolates, such as HCV1a, HCV1b, etc. A number of conserved and variable regions are known between these strains and, in general, the amino acid sequences of epitopes derived from these regions will have a high degree of sequence homology, e.g., amino acid sequence homology of more than 30%, preferably more than 40%, when the two sequences are aligned. Thus, for example, the term "NS4" 20 polypeptide refers to native NS4 from any of the various HCV strains, as well as NS4 analogs, muteins and immunogenic fragments, as defined further below.

By an "E1 polypeptide" is meant a molecule derived from an HCV E1 region. The mature E1 region of HCV-1 begins at approximately amino acid 192 of the polyprotein and continues to approximately amino acid 383, numbered relative to the full-length HCV-1 polyprotein. (See, Figures 1 and 3A-3C. Amino acids 192-383 of Figures 3A-3C correspond to amino acid positions 20-211 of SEQ ID NO:4.) Amino acids at around 173 through approximately 191 (amino acids 1-19 of SEQ ID NO: 4) serve as a signal sequence for E1. Thus, by an "E1 polypeptide" is meant either a precursor E1 protein, including the signal sequence, or a mature E1 polypeptide which lacks this sequence, or even an E1 polypeptide with a heterologous signal sequence.

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The E1 polypeptide includes a C-terminal membrane anchor sequence which occurs at approximately amino acid positions 360-383 (see, International Publication No. WO 96/04301, published February 15, 1996). An E1 polypeptide, as defined herein, may or may not include the C-terminal anchor sequence or portions thereof.

By an "E2 polypeptide" is meant a molecule derived from an HCV E2 region. The mature E2 region of HCV-1 begins at approximately amino acid 383-385. numbered relative to the full-length HCV-1 polyprotein. (See, Figures 1 and 3A-3C. Amino acids 383-385 of Figures 3A-3C correspond to amino acid positions 211-213 of SEQ ID NO:4.) A signal peptide begins at approximately amino acid 364 of the polyprotein. Thus, by an "E2 polypeptide" is meant either a precursor E2 protein. including the signal sequence, or a mature E2 polypeptide which lacks this sequence. or even an E2 polypeptide with a heterologous signal sequence. The E2 polypeptide includes a C-terminal membrane anchor sequence which occurs at approximately amino acid positions 715-730 and may extend as far as approximately amino acid residue 746 (see, Lin et al., J. Virol. (1994) 68:5063-5073). An E2 polypeptide, as defined herein, may or may not include the C-terminal anchor sequence or portions thereof. Moreover, an E2 polypeptide may also include all or a portion of the p7 region which occurs immediately adjacent to the C-terminus of E2. As shown in Figures 1 and 3A-3C, the p7 region is found at positions 747-809, numbered relative to the full-length HCV-1 polyprotein (amino acid positions 575-637 of SEO ID NO:4). Additionally, it is known that multiple species of HCV E2 exist (Spaete et al., Virol. (1992) 188:819-830; Selby et al., J. Virol. (1996) 70:5177-5182; Grakoui et al., J. Virol. (1993) 67:1385-1395; Tomei et al., J. Virol. (1993) 67:4017-4026). Accordingly, for purposes of the present invention, the term "E2" encompasses any of these species of E2 including, without limitation, species that have deletions of 1-20 or more of the amino acids from the N-terminus of the E2, such as, e.g, deletions of 1, 2, 3, 4, 5....10...15, 16, 17, 18, 19... etc. amino acids. Such E2 species include those beginning at amino acid 387, amino acid 402, amino acid 403, etc.

Representative E1 and E2 regions from HCV-1 are shown in Figures 3A-3C and SEQ ID NO:4. For purposes of the present invention, the E1 and E2 regions are defined with respect to the amino acid number of the polyprotein encoded by the

genome of HCV-1, with the initiator methionine being designated position 1. See, e.g., Choo et al., *Proc. Natl. Acad. Sci. USA* (1991) <u>88</u>:2451-2455. However, it should be noted that the term an "E1 polypeptide" or an "E2 polypeptide" as used herein is not limited to the HCV-1 sequence. In this regard, the corresponding E1 or E2 regions in other HCV isolates can be readily determined by aligning sequences from the isolates in a manner that brings the sequences into maximum alignment. This can be performed with any of a number of computer software packages, such as ALIGN 1.0, available from the University of Virginia, Department of Biochemistry (Attn: Dr. William R. Pearson). See, Pearson et al., *Proc. Natl. Acad. Sci. USA* (1988) <u>85</u>:2444-2448.

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Furthermore, an "E1 polypeptide" or an "E2 polypeptide" as defined herein is not limited to a polypeptide having the exact sequence depicted in the Figures.

Indeed, the HCV genome is in a state of constant flux *in vivo* and contains several variable domains which exhibit relatively high degrees of variability between isolates. A number of conserved and variable regions are known between these strains and, in general, the amino acid sequences of epitopes derived from these regions will have a high degree of sequence homology, e.g., amino acid sequence homology of more than 30%, preferably more than 40%, more than 60%, and even more than 80-90% homology, when the two sequences are aligned. It is readily apparent that the terms encompass E1 and E2 polypeptides from any of the various HCV strains and isolates including isolates having any of the 6 genotypes of HCV described in Simmonds et al., *J. Gen. Virol.* (1993) 74:2391-2399 (e.g., strains 1, 2, 3, 4 etc.), as well as newly identified isolates, and subtypes of these isolates, such as HCV1a, HCV1b etc.

Thus, for example, the term "E1" or "E2" polypeptide refers to native E1 or E2 sequences from any of the various HCV strains, as well as analogs, muteins and immunogenic fragments, as defined further below. The complete genotypes of many of these strains are known. See, e.g., U.S. Patent No. 6,150,087 and GenBank Accession Nos. AJ238800 and AJ238799.

Additionally, the terms "E1 polypeptide" and "E2 polypeptide" encompass proteins which include modifications to the native sequence, such as internal deletions, additions and substitutions (generally conservative in nature). These

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modifications may be deliberate, as through site-directed mutagenesis, or may be accidental, such as through naturally occurring mutational events. All of these modifications are encompassed in the present invention so long as the modified E1 and E2 polypeptides function for their intended purpose. Thus, for example, if the E1 and/or E2 polypeptides are to be used in vaccine compositions, the modifications must be such that immunological activity (i.e., the ability to elicit a humoral or cellular immune response to the polypeptide) is not lost.

By "E1E2" complex is meant a protein containing at least one E1 polypeptide and at least one E2 polypeptide, as described above. Such a complex may also include all or a portion of the p7 region which occurs immediately adjacent to the C-terminus of E2. As shown in Figures 1 and 3A-3C, the p7 region is found at positions 747-809, numbered relative to the full-length HCV-1 polyprotein (amino acid positions 575-637 of SEQ ID NO:4). A representative E1E2 complex which includes the p7 protein is termed "E1E2₈₀₉" herein.

The mode of association of E1 and E2 in an E1E2 complex is immaterial. The E1 and E2 polypeptides may be associated through non-covalent interactions such as through electrostatic forces, or by covalent bonds. For example, the E1E2 polypeptides of the present invention may be in the form of a fusion protein which includes an immunogenic E1 polypeptide and an immunogenic E2 polypeptide, as defined above. The fusion may be expressed from a polynucleotide encoding an E1E2 chimera. Alternatively, E1E2 complexes may form spontaneously simply by mixing E1 and E2 proteins which have been produced individually. Similarly, when coexpressed and secreted into media, the E1 and E2 proteins can form a complex spontaneously. Thus, the term encompasses E1E2 complexes (also called aggregates) that spontaneously form upon purification of E1 and/or E2. Such aggregates may include one or more E1 monomers in association with one or more E2 monomers. The number of E1 and E2 monomers present need not be equal so long as at least one E1 monomer and one E2 monomer are present. Detection of the presence of an E1E2 complex is readily determined using standard protein detection techniques such as polyacrylamide gel electrophoresis and immunological techniques such as immunoprecipitation.

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The terms "analog" and "mutein" refer to biologically active derivatives of the reference molecule, or fragments of such derivatives, that retain desired activity, such as the ability to stimulate a cell-mediated immune response, as defined below. In general, the term "analog" refers to compounds having a native polypeptide sequence and structure with one or more amino acid additions, substitutions (generally conservative in nature) and/or deletions, relative to the native molecule, so long as the modifications do not destroy immunogenic activity. The term "mutein" refers to peptides having one or more peptide mimics ("peptoids"), such as those described in International Publication No. WO 91/04282. Preferably, the analog or mutein has at least the same immunoactivity as the native molecule. Methods for making polypeptide analogs and muteins are known in the art and are described further below.

As explained above, analogs generally include substitutions that are conservative in nature, i.e., those substitutions that take place within a family of amino acids that are related in their side chains. Specifically, amino acids are generally divided into four families: (1) acidic -- aspartate and glutamate; (2) basic -- lysine, arginine, histidine; (3) non-polar – alanine, valine, leucine, isoleucine, proline, phenylalanine, methionine, tryptophan; and (4) uncharged polar -- glycine, asparagine, glutamine, cysteine, serine threonine, tyrosine. Phenylalanine, tryptophan, and tyrosine are sometimes classified as aromatic amino acids. For example, it is reasonably predictable that an isolated replacement of leucine with isoleucine or valine, an aspartate with a glutamate, a threonine with a serine, or a similar conservative replacement of an amino acid with a structurally related amino acid, will not have a major effect on the biological activity. For example, the polypeptide of interest may include up to about 5-10 conservative or non-conservative amino acid substitutions, or even up to about 15-25 conservative or non-conservative amino acid substitutions, or any integer between 5-25, so long as the desired function of the molecule remains intact. One of skill in the art may readily determine regions of the molecule of interest that can tolerate change by reference to Hopp/Woods and Kyte-Doolittle plots, well known in the art.

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By "modified NS3" is meant an NS3 polypeptide with a modification such that protease activity of the NS3 polypeptide is disrupted. The modification can include one or more amino acid additions, substitutions (generally non-conservative in nature) and/or deletions, relative to the native molecule, wherein the protease activity of the NS3 polypeptide is disrupted. Methods of measuring protease activity are discussed further below.

By "fragment" is intended a polypeptide consisting of only a part of the intact full-length polypeptide sequence and structure. The fragment can include a C-terminal deletion and/or an N-terminal deletion of the native polypeptide. An "immunogenic fragment" of a particular HCV protein will generally include at least about 5-10 contiguous amino acid residues of the full-length molecule, preferably at least about 15-25 contiguous amino acid residues of the full-length molecule, and most preferably at least about 20-50 or more contiguous amino acid residues of the full-length molecule, that define an epitope, or any integer between 5 amino acids and the full-length sequence, provided that the fragment in question retains immunogenic activity, as measured by the assays described herein.

The term "epitope" as used herein refers to a sequence of at least about 3 to 5, preferably about 5 to 10 or 15, and not more than about 1,000 amino acids (or any integer therebetween), which define a sequence that by itself or as part of a larger sequence, binds to an antibody generated in response to such sequence. There is no critical upper limit to the length of the fragment, which may comprise nearly the full-length of the protein sequence, or even a fusion protein comprising two or more epitopes from the HCV polyprotein. An epitope for use in the subject invention is not limited to a polypeptide having the exact sequence of the portion of the parent protein from which it is derived. Indeed, viral genomes are in a state of constant flux and contain several variable domains which exhibit relatively high degrees of variability between isolates. Thus the term "epitope" encompasses sequences identical to the native sequence, as well as modifications to the native sequence, such as deletions, additions and substitutions (generally conservative in nature).

Regions of a given polypeptide that include an epitope can be identified using any number of epitope mapping techniques, well known in the art. See, e.g., Epitope

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Mapping Protocols in Methods in Molecular Biology, Vol. 66 (Glenn E. Morris, Ed., 1996) Humana Press, Totowa, New Jersey. For example, linear epitopes may be determined by e.g., concurrently synthesizing large numbers of peptides on solid supports, the peptides corresponding to portions of the protein molecule, and reacting the peptides with antibodies while the peptides are still attached to the supports. Such techniques are known in the art and described in, e.g., U.S. Patent No. 4,708,871; Geysen et al. (1984) Proc. Natl. Acad. Sci. USA 81:3998-4002; Geysen et al. (1986) Molec. Immunol. 23:709-715. Similarly, conformational epitopes are readily identified by determining spatial conformation of amino acids such as by, e.g., x-ray crystallography and 2-dimensional nuclear magnetic resonance. See, e.g., Epitope Mapping Protocols, supra. Antigenic regions of proteins can also be identified using standard antigenicity and hydropathy plots, such as those calculated using, e.g., the Omiga version 1.0 software program available from the Oxford Molecular Group. This computer program employs the Hopp/Woods method, Hopp et al., Proc. Natl. Acad. Sci USA (1981) 78:3824-3828 for determining antigenicity profiles, and the Kyte-Doolittle technique, Kyte et al., J. Mol. Biol. (1982) 157:105-132 for hydropathy plots.

For a description of various HCV epitopes, see, e.g., Chien et al., *Proc. Natl. Acad. Sci. USA* (1992) <u>89</u>:10011-10015; Chien et al., *J. Gastroent. Hepatol.* (1993) <u>8</u>:S33-39; Chien et al., International Publication No. WO 93/00365; Chien, D.Y., International Publication No. WO 94/01778; and U.S. Patent Nos. 6,280,927 and 6,150,087.

As used herein, the term "conformational epitope" refers to a portion of a full-length protein, or an analog or mutein thereof, having structural features native to the amino acid sequence encoding the epitope within the full-length natural protein.

Native structural features include, but are not limited to, glycosylation and three dimensional structure. Preferably, a conformational epitope is produced recombinantly and is expressed in a cell from which it is extractable under conditions which preserve its desired structural features, e.g. without denaturation of the epitope. Such cells include bacteria, yeast, insect, and mammalian cells. Expression and isolation of recombinant conformational epitopes from the HCV polyprotein are

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described in e.g., International Publication Nos. WO 96/04301, WO 94/01778, WO 95/33053, WO 92/08734.

As used herein the term "T-cell epitope" refers to a feature of a peptide structure which is capable of inducing T-cell immunity towards the peptide structure or an associated hapten. T-cell epitopes generally comprise linear peptide determinants that assume extended conformations within the peptide-binding cleft of MHC molecules, (Unanue et al., Science (1987) 236:551-557). Conversion of polypeptides to MHC class II-associated linear peptide determinants (generally between 5-14 amino acids in length) is termed "antigen processing" which is carried out by antigen presenting cells (APCs). More particularly, a T-cell epitope is defined by local features of a short peptide structure, such as primary amino acid sequence properties involving charge and hydrophobicity, and certain types of secondary structure, such as helicity, that do not depend on the folding of the entire polypeptide. Further, it is believed that short peptides capable of recognition by helper T-cells are generally amphipathic structures comprising a hydrophobic side (for interaction with the MHC molecule) and a hydrophilic side (for interacting with the T-cell receptor). (Margalit et al., Computer Prediction of T-cell Epitopes, New Generation Vaccines Marcel-Dekker, Inc, ed. G.C. Woodrow et al., (1990) pp. 109-116) and further that the amphipathic structures have an α-helical configuration (see, e.g., Spouge et al., J. Immunol. (1987) 138:204-212; Berkower et al., J. Immunol. (1986) 136:2498-2503).

Hence, segments of proteins that include T-cell epitopes can be readily predicted using numerous computer programs. (See e.g., Margalit et al., Computer Prediction of T-cell Epitopes, New Generation Vaccines Marcel-Dekker, Inc, ed. G.C. Woodrow et al., (1990) pp. 109-116). Such programs generally compare the amino acid sequence of a peptide to sequences known to induce a T-cell response, and search for patterns of amino acids which are believed to be required for a T-cell epitope.

An "immunological response" to an HCV antigen (including both polypeptide and polynucleotides encoding polypeptides that are expressed *in vivo*) or composition is the development in a subject of a humoral and/or a cellular immune response to molecules present in the composition of interest. For purposes of the present invention, a "humoral immune response" refers to an immune response mediated by

antibody molecules, while a "cellular immune response" is one mediated by T-lymphocytes and/or other white blood cells. One important aspect of cellular immunity involves an antigen-specific response by cytolytic T-cells ("CTLs"). CTLs have specificity for peptide antigens that are presented in association with proteins encoded by the major histocompatibility complex (MHC) and expressed on the surfaces of cells. CTLs help induce and promote the intracellular destruction of intracellular microbes, or the lysis of cells infected with such microbes. Another aspect of cellular immunity involves an antigen-specific response by helper T-cells. Helper T-cells act to help stimulate the function, and focus the activity of, nonspecific effector cells against cells displaying peptide antigens in association with MHC molecules on their surface. A "cellular immune response" also refers to the production of cytokines, chemokines and other such molecules produced by activated T-cells and/or other white blood cells, including those derived from CD4+ and CD8+ T-cells.

A composition or vaccine that elicits a cellular immune response may serve to sensitize a vertebrate subject by the presentation of antigen in association with MHC molecules at the cell surface. The cell-mediated immune response is directed at, or near, cells presenting antigen at their surface. In addition, antigen-specific T-lymphocytes can be generated to allow for the future protection of an immunized host.

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The ability of a particular antigen to stimulate a cell-mediated immunological response may be determined by a number of assays, such as by lymphoproliferation (lymphocyte activation) assays, CTL cytotoxic cell assays, or by assaying for T-lymphocytes specific for the antigen in a sensitized subject. Such assays are well known in the art. See, e.g., Erickson et al., *J. Immunol.* (1993) 151:4189-4199; Doe et al., *Eur. J. Immunol.* (1994) 24:2369-2376; and the examples below.

Thus, an immunological response as used herein may be one which stimulates the production of CTLs, and/or the production or activation of helper T- cells. The antigen of interest may also elicit an antibody-mediated immune response. Hence, an immunological response may include one or more of the following effects: the production of antibodies by B-cells; and/or the activation of suppressor T-cells and/or

 $\gamma\delta$ T-cells directed specifically to an antigen or antigens present in the composition or vaccine of interest. These responses may serve to neutralize infectivity, and/or mediate antibody-complement, or antibody dependent cell cytotoxicity (ADCC) to provide protection or alleviation of symptoms to an immunized host. Such responses can be determined using standard immunoassays and neutralization assays, well known in the art.

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By "equivalent antigenic determinant" is meant an antigenic determinant from different sub-species or strains of HCV, such as from strains 1, 2, 3, etc., of HCV which antigenic determinants are not necessarily identical due to sequence variation, but which occur in equivalent positions in the HCV sequence in question. In general the amino acid sequences of equivalent antigenic determinants will have a high degree of sequence homology, e.g., amino acid sequence homology of more than 30%, usually more than 40%, such as more than 60%, and even more than 80-90% homology, when the two sequences are aligned.

A "coding sequence" or a sequence which "encodes" a selected polypeptide, is a nucleic acid molecule which is transcribed (in the case of DNA) and translated (in the case of mRNA) into a polypeptide *in vitro* or *in vivo* when placed under the control of appropriate regulatory sequences. The boundaries of the coding sequence are determined by a start codon at the 5' (amino) terminus and a translation stop codon at the 3' (carboxy) terminus. A transcription termination sequence may be located 3' to the coding sequence.

A "nucleic acid" molecule or "polynucleotide" can include both double- and single-stranded sequences and refers to, but is not limited to, cDNA from viral, procaryotic or eucaryotic mRNA, genomic DNA sequences from viral (e.g. DNA viruses and retroviruses) or procaryotic DNA, and especially synthetic DNA sequences. The term also captures sequences that include any of the known base analogs of DNA and RNA.

An "HCV polynucleotide" is a polynucleotide that encodes an HCV polypeptide, as defined above.

"Operably linked" refers to an arrangement of elements wherein the components so described are configured so as to perform their desired function. Thus,

a given promoter operably linked to a coding sequence is capable of effecting the expression of the coding sequence when the proper transcription factors, etc., are present. The promoter need not be contiguous with the coding sequence, so long as it functions to direct the expression thereof. Thus, for example, intervening untranslated yet transcribed sequences can be present between the promoter sequence and the coding sequence, as can transcribed introns, and the promoter sequence can still be considered "operably linked" to the coding sequence.

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"Recombinant" as used herein to describe a nucleic acid molecule means a polynucleotide of genomic, cDNA, viral, semisynthetic, or synthetic origin which, by virtue of its origin or manipulation is not associated with all or a portion of the polynucleotide with which it is associated in nature. The term "recombinant" as used with respect to a protein or polypeptide means a polypeptide produced by expression of a recombinant polynucleotide. In general, the gene of interest is cloned and then expressed in transformed organisms, as described further below. The host organism expresses the foreign gene to produce the protein under expression conditions.

A "control element" refers to a polynucleotide sequence which aids in the expression of a coding sequence to which it is linked. The term includes promoters, transcription termination sequences, upstream regulatory domains, polyadenylation signals, untranslated regions, including 5'-UTRs and 3'-UTRs and when appropriate, leader sequences and enhancers, which collectively provide for the transcription and translation of a coding sequence in a host cell.

A "promoter" as used herein is a DNA regulatory region capable of binding RNA polymerase in a host cell and initiating transcription of a downstream (3' direction) coding sequence operably linked thereto. For purposes of the present invention, a promoter sequence includes the minimum number of bases or elements necessary to initiate transcription of a gene of interest at levels detectable above background. Within the promoter sequence is a transcription initiation site, as well as protein binding domains (consensus sequences) responsible for the binding of RNA polymerase. Eucaryotic promoters will often, but not always, contain "TATA" boxes and "CAT" boxes.

A control sequence "directs the transcription" of a coding sequence in a cell when RNA polymerase will bind the promoter sequence and transcribe the coding sequence into mRNA, which is then translated into the polypeptide encoded by the coding sequence.

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"Expression cassette" or "expression construct" refers to an assembly which is capable of directing the expression of the sequence(s) or gene(s) of interest. The expression cassette includes control elements, as described above, such as a promoter which is operably linked to (so as to direct transcription of) the sequence(s) or gene(s) of interest, and often includes a polyadenylation sequence as well. Within certain embodiments of the invention, the expression cassette described herein may be contained within a plasmid construct. In addition to the components of the expression cassette, the plasmid construct may also include, one or more selectable markers, a signal which allows the plasmid construct to exist as single-stranded DNA (e.g., a M13 origin of replication), at least one multiple cloning site, and a "mammalian" origin of replication (e.g., a SV40 or adenovirus origin of replication).

"Transformation," as used herein, refers to the insertion of an exogenous polynucleotide into a host cell, irrespective of the method used for insertion: for example, transformation by direct uptake, transfection, infection, and the like. For particular methods of transfection, see further below. The exogenous polynucleotide may be maintained as a nonintegrated vector, for example, an episome, or alternatively, may be integrated into the host genome.

A "host cell" is a cell which has been transformed, or is capable of transformation, by an exogenous DNA sequence.

By "isolated" is meant, when referring to a polypeptide, that the indicated molecule is separate and discrete from the whole organism with which the molecule is found in nature or is present in the substantial absence of other biological macromolecules of the same type. The term "isolated" with respect to a polynucleotide is a nucleic acid molecule devoid, in whole or part, of sequences normally associated with it in nature; or a sequence, as it exists in nature, but having heterologous sequences in association therewith; or a molecule disassociated from the chromosome.

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The term "purified" as used herein preferably means at least 75% by weight, more preferably at least 85% by weight, more preferably still at least 95% by weight, and most preferably at least 98% by weight, of biological macromolecules of the same type are present.

"Homology" refers to the percent identity between two polynucleotide or two polypeptide moieties. Two DNA, or two polypeptide sequences are "substantially homologous" to each other when the sequences exhibit at least about 50%, preferably at least about 75%, more preferably at least about 80%-85%, preferably at least about 90%, and most preferably at least about 95%-98%, or more, sequence identity over a defined length of the molecules. As used herein, substantially homologous also refers to sequences showing complete identity to the specified DNA or polypeptide sequence.

In general, "identity" refers to an exact nucleotide-to-nucleotide or amino acidto-amino acid correspondence of two polynucleotides or polypeptide sequences, respectively. Percent identity can be determined by a direct comparison of the sequence information between two molecules by aligning the sequences, counting the exact number of matches between the two aligned sequences, dividing by the length of the shorter sequence, and multiplying the result by 100. Readily available computer programs can be used to aid in the analysis, such as ALIGN, Dayhoff, M.O. in Atlas of Protein Sequence and Structure M.O. Dayhoff ed., 5 Suppl. 3:353-358, National biomedical Research Foundation, Washington, DC, which adapts the local homology algorithm of Smith and Waterman Advances in Appl. Math. 2:482-489, 1981 for peptide analysis. Programs for determining nucleotide sequence identity are available in the Wisconsin Sequence Analysis Package, Version 8 (available from Genetics Computer Group, Madison, WI) for example, the BESTFIT, FASTA and GAP programs, which also rely on the Smith and Waterman algorithm. These programs are readily utilized with the default parameters recommended by the manufacturer and described in the Wisconsin Sequence Analysis Package referred to above. For example, percent identity of a particular nucleotide sequence to a reference sequence can be determined using the homology algorithm of Smith and Waterman with a default scoring table and a gap penalty of six nucleotide positions.

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Another method of establishing percent identity in the context of the present invention is to use the MPSRCH package of programs copyrighted by the University of Edinburgh, developed by John F. Collins and Shane S. Sturrok, and distributed by IntelliGenetics, Inc. (Mountain View, CA). From this suite of packages the Smith-Waterman algorithm can be employed where default parameters are used for the scoring table (for example, gap open penalty of 12, gap extension penalty of one, and a gap of six). From the data generated the "Match" value reflects "sequence identity." Other suitable programs for calculating the percent identity or similarity between sequences are generally known in the art, for example, another alignment program is BLAST, used with default parameters. For example, BLASTN and BLASTP can be used using the following default parameters: genetic code = standard; filter = none; strand = both; cutoff = 60; expect = 10; Matrix = BLOSUM62; Descriptions = 50 sequences; sort by = HIGH SCORE; Databases = non-redundant, GenBank + EMBL + DDBJ + PDB + GenBank CDS translations + Swiss protein + Spupdate + PIR. Details of these programs can be found at the following internet address: http://www.ncbi.nlm.gov/cgi-bin/BLAST.

Alternatively, homology can be determined by hybridization of polynucleotides under conditions which form stable duplexes between homologous regions, followed by digestion with single-stranded-specific nuclease(s), and size determination of the digested fragments. DNA sequences that are substantially homologous can be identified in a Southern hybridization experiment under, for example, stringent conditions, as defined for that particular system. Defining appropriate hybridization conditions is within the skill of the art. See, e.g., Sambrook et al., supra; DNA Cloning, supra; Nucleic Acid Hybridization, supra.

By "nucleic acid immunization" is meant the introduction of a nucleic acid molecule encoding one or more selected antigens into a host cell, for the *in vivo* expression of the antigen or antigens. The hucleic acid molecule can be introduced directly into the recipient subject, such as by injection, inhalation, oral, intranasal and mucosal administration, or the like, or can be introduced *ex vivo*, into cells which have been removed from the host. In the latter case, the transformed cells are reintroduced

into the subject where an immune response can be mounted against the antigen encoded by the nucleic acid molecule.

As used herein, "treatment" refers to any of (i) the prevention of infection or reinfection, as in a traditional vaccine, (ii) the reduction or elimination of symptoms, and (iii) the substantial or complete elimination of the pathogen in question.

Treatment may be effected prophylactically (prior to infection) or therapeutically (following infection).

By "vertebrate subject" is meant any member of the subphylum cordata, including, without limitation, humans and other primates, including non-human primates such as chimpanzees and other apes and monkey species; farm animals such as cattle, sheep, pigs, goats and horses; domestic mammals such as dogs and cats; laboratory animals including rodents such as mice, rats and guinea pigs; birds, including domestic, wild and game birds such as chickens, turkeys and other gallinaceous birds, ducks, geese, and the like. The term does not denote a particular age. Thus, both adult and newborn individuals are intended to be covered. The invention described herein is intended for use in any of the above vertebrate species, since the immune systems of all of these vertebrates operate similarly.

II. Modes of Carrying out the Invention

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Before describing the present invention in detail, it is to be understood that this invention is not limited to particular formulations or process parameters as such may, of course, vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments of the invention only, and is not intended to be limiting.

Although a number of compositions and methods similar or equivalent to those described herein can be used in the practice of the present invention, the preferred materials and methods are described herein.

It is a discovery of the present invention that fusion proteins, combinations of the individual components of these fusions, and polynucleotides encoding the same, comprising an NS3, an NS4, and an NS5a polypeptide with or without a core polypeptide, or an NS3, an NS4, an NS5a, and an NS5b polypeptide, with or without a

core polypeptide, of an HCV virus can be used to activate HCV-specific T cells, i.e., T cells which recognize epitopes of these polypeptides.

The present invention also pertains to compositions comprising HCV nonstructural fusion proteins and HCV E1E2 complexes, as well as compositions comprising polynucleotides encoding the same or combinations of polypeptides and polynucleotides.

The proteins, polynucleotides, compositions and combinations of the present invention can be used to stimulate a cellular immune response, such as to activate HCV-specific T cells, i.e., T cells which recognize epitopes of these polypeptides. Activation of HCV-specific T cells provides both *in vitro* and *in vivo* model systems for the development of HCV vaccines, particularly for identifying HCV polypeptide epitopes associated with a response. The compositions can also be used to generate an immune response against HCV in a mammal, particularly a CTL response for either therapeutic or prophylactic purposes.

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Fusion Proteins

The genomes of HCV strains contain a single open reading frame of approximately 9,000 to 12,000 nucleotides, which is transcribed into a polyprotein. As shown in Figure 1 and the table below, an HCV polyprotein, upon cleavage, produces at least ten distinct products, in the order of NH₂- Core-E1-E2-p7-NS2-NS3-NS4a-NS4b-NS5a-NS5b-COOH. The core polypeptide occurs at positions 1-191, numbered relative to HCV-1 (see, Choo et al. (1991) *Proc. Natl. Acad. Sci. USA* 88:2451-2455, for the HCV-1 genome). This polypeptide is further processed to produce an HCV polypeptide with approximately amino acids 1-173. The envelope polypeptides, E1 and E2, occur at about positions 192-383 and 384-746, respectively. The P7 domain is found at about positions 747-809. NS2 is an integral membrane protein with proteolytic activity and is found at about positions 810-1026 of the polyprotein. NS2, in combination with NS3, (found at about positions 1027-1657), cleaves the NS2-NS3 sissle bond which in turn generates the NS3 N-terminus and releases a large polyprotein that includes both serine protease and RNA helicase activities. The NS3 protease, found at about positions 1027-1207, serves to process

the remaining polyprotein. The helicase activity is found at about positions 1193-1657. NS3 liberates an NS3 cofactor (NS4a, found about positions 1658-1711), two proteins (NS4b found at about positions 1712-1972, and NS5a found at about positions 1973-2420), and an RNA-dependent RNA polymerase (NS5b found at about positions 2421-3011). Completion of polyprotein maturation is initiated by autocatalytic cleavage at the NS3-Ns4a junction, catalyzed by the NS3 serine protease.

Approximate Boundaries*

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- (0000)	2, 22 2
E1	192-383
E2	384-746
P7	747-809
NS2	810-1026
NS3	1027-1657
NS4a	1658-1711
NS4b	1712-1972
NS5a	1973-2420

Domain

C (core)

NS5b

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*Numbered relative to HCV-1. See, Choo et al. (1991) *Proc. Natl. Acad. Sci. USA* 88:2451-2455.

2421-3011

Fusion proteins for use in the compositions and methods, and polynucleotides encoding therefor, include or encode an NS3 polypeptide, an NS4 (NS4a and/or NS4b) polypeptide, an NS5a polypeptide and, optionally, an NS5b polypeptide. The fusion proteins may or may not include all or part of the core region. In certain embodiments, none of the core region is present in the compositions. The nonstructural regions need not be in the order in which they naturally occur in the native HCV polyprotein. Thus, for example, the NS5b polypeptide may be at the N–and/or C-terminus of the fusion or may be found internally. These polypeptides may

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be derived from the same HCV isolate, or from different strains and isolates including isolates having any of the various HCV genotypes, to provide increased protection against a broad range of HCV genotypes. Additionally, polypeptides can be selected based on the particular viral clades endemic in specific geographic regions where vaccine compositions containing the fusions will be used. It is readily apparent that the subject fusions provide an effective means of treating HCV infection in a wide variety of contexts.

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In one embodiment, the fusion protein of the present invention includes an NS3 polypeptide that has been modified to inhibit protease activity, such that further cleavage of the fusion is inhibited. The NS3 polypeptide can be modified by deletion of all or a portion of the NS3 protease domain. Alternatively, proteolytic activity can be inhibited by substitutions of amino acids within active regions of the protease domain. Finally, additions of amino acids to active regions of the domain, such that the catalytic site is modified, will also serve to inhibit proteolytic activity.

As explained above, the protease activity is found at about amino acid positions 1027-1207, numbered relative to the full-length HCV-1 polyprotein (see, Choo et al., *Proc. Natl. Acad. Sci. USA* (1991) 88:2451-2455), positions 2-182 of Figure 3. The structure of the NS3 protease and active site are known. See, e.g., De Francesco et al., *Antivir. Ther.* (1998) 3:99-109; Koch et al., *Biochemistry* (2001) 40:631-640. Thus, deletions or modifications to the native sequence will typically occur at or near the active site of the molecule. Particularly, it is desirable to modify or make deletions to one or more amino acids occurring at positions 1- or 2-182, preferably 1- or 2-170, or 1- or 2-155 of Figure 3. Preferred modifications are to the catalytic triad at the active site of the protease, i.e., H, D or S residues, in order to inactivate the protease. These residues occur at positions 1083, 1105 and 1165, respectively, numbered relative to the full-length HCV polyprotein (positions 58, 80 and 140, respectively, of Figure 3). Such modifications will suppress proteolytic cleavage while maintaining T-cell epitopes.

One of skill in the art can readily determine portions of the NS3 protease to delete in order to disrupt activity. The presence or absence of activity can be determined using methods known to those of skill in the art.

For example, protease activity or lack thereof may be determined using assays well known in the art. See, e.g., Takeshita et al., Anal. Biochem. (1997) 247:242-246; Kakiuchi et al., J. Biochem. (1997) 122:749-755; Sali et al., Biochemistry (1998) 37:3392-3401; Cho et al., J. Virol. Meth. (1998) 72:109-115; Cerretani et al., Anal. Biochem. (1999) 266:192-197; Zhang et al., Anal. Biochem. (1999) 270:268-275; Kakiuchi et al., J. Virol. Meth. (1999) 80:77-84; Fowler et al., J. Biomol. Screen. (2000) 5:153-158; and Kim et al., Anal. Biochem. (2000) 284:42-48.

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The NS3, NS4, NS5a, and NS5b polypeptides present in the various fusions described above can either be full-length polypeptides or portions of NS3, NS4 (NS4a and/or NS4b), NS5a, and NS5b polypeptides. The portions of NS3, NS4, NS5a, and NS5b polypeptides making up the fusion protein preferably comprise at least one epitope, which is recognized by a T cell receptor on an activated T cell, such as 2152-HEYPVGSQL-2160 (SEQ ID NO:1) and/or 2224-AELIEANLLWRQEMG-2238 (SEQ ID NO:2). Epitopes of NS3, NS4 (NS4a and NS4b), NS5a, NS5b, NS3NS4NS5a, and NS3NS4NS5aNS5b can be identified by several methods. For example, NS3, NS4, NS5a, NS5b polypeptides or fusion proteins comprising any combination of the above, can be isolated, for example, by immunoaffinity purification using a monoclonal antibody for the polypeptide or protein. The isolated protein sequence can then be screened by preparing a series of short peptides by proteolytic cleavage of the purified protein, which together span the entire protein sequence. By starting with, for example, 100-mer polypeptides, each polypeptide can be tested for the presence of epitopes recognized by a T-cell receptor on an HCV-activated T cell, progressively smaller and overlapping fragments can then be tested from an identified 100-mer to map the epitope of interest.

Epitopes recognized by a T-cell receptor on an HCV-activated T cell can be identified by, for example, ⁵¹Cr release assay or by lymphoproliferation assay (see the examples). In a ⁵¹Cr release assay, target cells can be constructed that display the epitope of interest by cloning a polynucleotide encoding the epitope into an expression vector and transforming the expression vector into the target cells. HCV-specific CD8⁺ T cells will lyse target cells displaying, for example, an NS3, NS4, NS5a, NS5b, NS3NS4NS5a, or NS3NS4NS5aNS5b epitope and will not lyse cells

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that do not display such an epitope. In a lymphoproliferation assay, HCV-activated CD4⁺ T cells will proliferate when cultured with, for example, an NS3, NS4, NS5a, NS5b, NS3NS4NS5a, or NS3NS4NS5aNS5b epitopic peptide, but not in the absence of an HCV epitopic peptide.

NS3, NS4, NS5a, and NS5b polypeptides can occur in any order in the fusion protein. If desired, at least 2, 3, 4, 5, 6, 7, 8, 9, or 10 or more of one or more of the polypeptides may occur in the fusion protein. Multiple viral strains of HCV occur, and NS3, NS4, NS5a, and NS5b polypeptides of any of these strains can be used in a fusion protein. A representative fusion protein for use in the present invention is shown if Figures 5A-5J. The depicted sequence includes amino acids 1242-3011 of the HCV polyprotein (representing polypeptides from NS3, NS4, NS5a and NS5b) with amino acids 1-121 of the HCV polyprotein (representing a polypeptide from the core region) fused to the C-terminus of NS5b. This numbering is relative to the HCV-1 polyprotein.

Nucleic acid and amino acid sequences of a number of HCV strains and isolates, including nucleic acid and amino acid sequences of NS3, NS4, NS5a, NS5b genes and polypeptides have been determined. For example, isolate HCV J1.1 is described in Kubo et al. (1989) Japan. Nucl. Acids Res. 17:10367-10372; Takeuchi et al. (1990) Gene 91:287-291; Takeuchi et al. (1990) J. Gen. Virol. 71:3027-3033; and Takeuchi et al. (1990) Nucl. Acids Res. 18:4626. The complete coding sequences of two independent isolates, HCV-J and BK, are described by Kato et al., (1990) Proc. Natl. Acad. Sci. USA 87:9524-9528 and Takamizawa et al., (1991) J. Virol. 65:1105-1113 respectively.

Publications that describe HCV-1 isolates include Choo et al. (1990) Brit. Med. Bull. 46:423-441; Choo et al. (1991) Proc. Natl. Acad. Sci. USA 88:2451-2455 and Han et al. (1991) Proc. Natl. Acad. Sci. USA 88:1711-1715. HCV isolates HC-J1 and HC-J4 are described in Okamoto et al. (1991) Japan J. Exp. Med. 60:167-177. HCV isolates HCT 18~, HCT 23, Th, HCT 27, EC1 and EC10 are described in Weiner et al. (1991) Virol. 180:842-848. HCV isolates Pt-1, HCV-K1 and HCV-K2 are described in Enomoto et al. (1990) Biochem. Biophys. Res.

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Commun. 170:1021-1025. HCV isolates A, C, D & E are described in Tsukiyama-Kohara et al. (1991) Virus Genes 5:243-254.

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Each of the NS3, NS4, NS5a, and NS5b components of a fusion protein can be obtained from the same HCV strain or isolate or from different HCV strains or isolates. Fusion proteins comprising HCV polypeptides from, for example, the NS3 polypeptide can be derived from a first strain of HCV, and the NS4, and NS5a polypeptides can be derived from a second strain of HCV. Alternatively, the NS4 polypeptide can be derived from a first strain of HCV, and the NS3 and NS5a polypeptides can be derived from a second strain of HCV. Optionally, the NS5a polypeptide can be derived from a first strain of HCV, and the NS3 and NS4 polypeptides can be derived from a second strain of HCV. NS3, NS4 and NS5a polypeptides can be derived from different HCV strains can also be used in an HCV fusion protein. Similarly, in a fusion protein comprising NS5b, at least one of the NS3, NS4, NS5a, and NS5b polypeptides can be derived from a different HCV strain than the other polypeptides. Optionally, NS3, NS4, NS5a, and NS5b polypeptides that are each derived from different HCV strains can also be used in an NS3NS4NS5aNS5b fusion protein.

In addition to NS3, NS4a, NS4b, NS5a and NS5b, the fusion proteins can contain other polypeptides derived from the HCV polyprotein. For example, it may be desirable to include polypeptides derived from the core region of the HCV polyprotein. This region occurs at amino acid positions 1-191 of the HCV polyprotein, numbered relative to HCV-1. Either the full-length protein, fragments thereof, such as amino acids 1-150, e.g., amino acids 1-130, 1-120, for example, amino acids 1-121, 1-122, 1-123, etc., or smaller fragments containing epitopes of the full-length protein may be used in the subject fusions, such as those epitopes found between amino acids 10-53, amino acids 10-45, amino acids 67-88, amino acids 120-130, or any of the core epitopes identified in, e.g., Houghton et al., U.S. Patent No. 5,350,671; Chien et al., *Proc. Natl. Acad. Sci. USA* (1992) 89:10011-10015; Chien et al., *J. Gastroent. Hepatol.* (1993) 8:S33-39; Chien et al., International Publication No. WO 93/00365; Chien, D.Y., International Publication No. WO 94/01778; and U.S. Patent Nos. 6,280,927 and 6,150,087. Moreover, a

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protein resulting from a frameshift in the core region of the polyprotein, such as described in International Publication No. WO 99/63941, may be used. The fusions may also contain polynucleotides encoding E1E2 polypeptides, as described further below.

Preferably, the above-described fusion proteins, as well as the individual components of these proteins, are produced recombinantly. A polynucleotide encoding these proteins can be introduced into an expression vector which can be expressed in a suitable expression system. A variety of bacterial, yeast, mammalian and insect expression systems are available in the art and any such expression system can be used. Optionally, a polynucleotide encoding these proteins can be translated in a cell-free translation system. Such methods are well known in the art. The proteins also can be constructed by solid phase protein synthesis.

If desired, the fusion proteins, or the individual components of these proteins, also can contain other non-HCV amino acid sequences, such as amino acid linkers or signal sequences, as well as ligands useful in protein purification, such as glutathione-S-transferase and staphylococcal protein A.

E1E2 Polypeptides

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As explained above, the compositions of the present invention may also include E1 and E2 polypeptides, complexes of these polypeptides or polynucleotides encoding the same. The E1 and E2 polypeptides and complexes thereof can be provided independent of the nonstructural fusion protein or can be incorporated into the same fusion. Moreover, E1E2 complexes can be provided as proteins, or as polynucleotides encoding the same.

In this regard, E1, E2 and p7 are known to contain human T-cell epitopes (both CD4+ and CD8+) and including one or more of these epitopes serves to increase vaccine efficacy as well as to increase protective levels against multiple HCV genotypes. Moreover, multiple copies of specific, conserved T-cell epitopes can also be used in E1E2 complexes, such as a composite of epitopes from different genotypes.

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As explained above, the E1 and E2 polypeptides that make up the E1E2 complexes can be associated either through non-covalent or covalent interactions. Such complexes may be made up of immunogenic fragments of E1 and E2 which comprise epitopes. For example, fragments of E1 polypeptides can comprise from about 5 to nearly the full-length of the molecule, such as 6, 10, 25, 50, 75, 100, 125, 150, 175, 185 or more amino acids of an E1 polypeptide, or any integer between the stated numbers. Similarly, fragments of E2 polypeptides can comprise 6, 10, 25, 50, 75, 100, 150, 200, 250, 300, or 350 amino acids of an E2 polypeptide, or any integer between the stated numbers. The E1 and E2 polypeptides may be from the same or different HCV strains. For example, epitopes derived from, e.g., the hypervariable region of E2, such as a region spanning amino acids 384-410 or 390-410, can be included in the E2 polypeptide. A particularly effective E2 epitope to incorporate into the E2 sequence or E1E2 complexes is one which includes a consensus sequence derived from this region, such as the consensus sequence Gly-Ser-Ala-Ala-Arg-Thr-Thr-Ser-Gly-Phe-Val-Ser-Leu-Phe-Ala-Pro-Gly-Ala-Lys-Gln-Asn (SEQ ID NO:5), which represents a consensus sequence for amino acids 390-410 of the HCV type 1 genome. Additional epitopes of E1 and E2 are known and described in, e.g., Chien et al., International Publication No. WO 93/00365.

Moreover, the E1 and E2 polypeptides may lack all or a portion of the membrane spanning domain. The membrane anchor sequence functions to associate the polypeptide to the endoplasmic reticulum. Normally, such polypeptides are capable of secretion into growth medium in which an organism expressing the protein is cultured. However, as described in International Publication No. WO 98/50556, such polypeptides may also be recovered intracellularly. Secretion into growth medium is readily determined using a number of detection techniques, including, e.g., polyacrylamide gel electrophoresis and the like, and immunological techniques such as immunoprecipitation assays as described in, e.g., International Publication No. WO 96/04301, published February 15, 1996. With E1, generally polypeptides terminating with about amino acid position 370 and higher (based on the numbering of HCV1 E1) will be retained by the ER and hence not secreted into growth media. With E2, polypeptides terminating with about amino acid position

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731 and higher (also based on the numbering of the HCV1 E2 sequence) will be retained by the ER and not secreted. (See, e.g., International Publication No. WO 96/04301, published February 15, 1996). It should be noted that these amino acid positions are not absolute and may vary to some degree. Thus, the present invention contemplates the use of E1 and E2 polypeptides which retain the transmembrane binding domain, as well as polypeptides which lack all or a portion of the transmembrane binding domain, including E1 polypeptides terminating at about amino acids 369 and lower, and E2 polypeptides, terminating at about amino acids 730 and lower, are intended to be captured by the present invention. Furthermore, the C-terminal truncation can extend beyond the transmembrane spanning domain towards the N-terminus. Thus, for example, E1 truncations occurring at positions lower than, e.g., 360 and E2 truncations occurring at positions lower than, e.g., 715, are also encompassed by the present invention. All that is necessary is that the truncated E1 and E2 polypeptides remain functional for their intended purpose. However, particularly preferred truncated E1 constructs are those that do not extend beyond about amino acid 300. Most preferred are those terminating at position 360. Preferred truncated E2 constructs are those with C-terminal truncations that do not extend beyond about amino acid position 715. Particularly preferred E2 truncations are those molecules truncated after any of amino acids 715-730, such as 725. If truncated molecules are used, it is preferable to use E1 and E2 molecules that are both truncated.

E2 exists as multiple species (Spaete et al., Virol. (1992) 188:819-830; Selby et al., J. Virol. (1996) 70:5177-5182; Grakoui et al., J. Virol. (1993) 67:1385-1395; Tomei et al., J. Virol. (1993) 67:4017-4026) and clipping and proteolysis may occur at the N- and C-termini of the E1 and E2 polypeptides. Thus, an E2 polypeptide for use herein may comprise at least amino acids 405-661, e.g., 400, 401, 402... to 661, such as 384-661, 384-715, 384-746, 384-749 or 384-809, or 384 to any C-terminus between 661-809, of an HCV polyprotein, numbered relative to the full-length HCV-1 polyprotein. Similarly, preferable E1 polypeptides for use herein can comprise amino acids 192-326, 192-330, 192-333, 192-360, 192-363, 192-383, or 192 to any C-terminus between 326-383, of an HCV polyprotein.

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E1 is fused to the desired portion of E2.

The E1 and E2 polypeptides and complexes thereof may also be present as asialoglycoproteins. Such asialoglycoproteins are produced by methods known in the art, such as by using cells in which terminal glycosylation is blocked. When these proteins are expressed in such cells and isolated by GNA lectin affinity chromatography, the E1 and E2 proteins aggregate spontaneously. Detailed methods for producing these E1E2 aggregates are described in, e.g., U.S. Patent No. 6,074,852. For example, E1E2 complexes are readily produced recombinantly, either as fusion proteins or by e.g., co-transfecting host cells with constructs encoding for the E1 and E2 polypeptides of interest. Co-transfection can be accomplished either in trans or cis, i.e., by using separate vectors or by using a single vector which bears both of the E1 and E2 genes. If done using a single vector, both genes can be driven by a single set of control elements or, alternatively, the genes can be present on the vector in individual expression cassettes, driven by individual control elements. Following expression, the E1 and E2 proteins will spontaneously associate. Alternatively, the complexes can be formed by mixing the individual proteins together which have been produced separately, either in purified or semi-purified form, or even by mixing culture media in which host cells expressing the proteins, have been cultured, if the proteins are secreted. Finally, the E1E2 complexes of the present invention may be expressed as a fusion protein wherein the desired portion of

Moreover, the E1E2 complexes may be present as a heterogeneous mixture of molecules, due to clipping and proteolytic cleavage, as described above. Thus, a composition including E1E2 complexes may include multiple species of E1E2, such as E1E2 terminating at amino acid 746 (E1E2₇₄₆), E1E2 terminating at amino acid 809 (E1E2₈₀₉), or any of the other various E1 and E2 molecules described above, such as E2 molecules with N-terminal truncations of from 1-20 amino acids, such as E2 species beginning at amino acid 387, amino acid 402, amino acid 403, etc.

E1E2 complexes are readily produced recombinantly, either as fusion proteins or by e.g., co-transfecting host cells with constructs encoding for the E1 and E2 polypeptides of interest. Co-transfection can be accomplished either in *trans* or *cis*, i.e., by using separate vectors or by using a single vector which bears both of the E1

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and E2 genes. If done using a single vector, both genes can be driven by a single set of control elements or, alternatively, the genes can be present on the vector in individual expression cassettes, driven by individual control elements. Following expression, the E1 and E2 proteins will spontaneously associate. Alternatively, the complexes can be formed by mixing the individual proteins together which have been produced separately, either in purified or semi-purified form, or even by mixing culture media in which host cells expressing the proteins, have been cultured, if the proteins are secreted. Finally, the E1E2 complexes of the present invention may be expressed as a fusion protein wherein the desired portion of E1 is fused to the desired portion of E2.

Methods for producing E1E2 complexes from full-length, truncated E1 and E2 proteins which are secreted into media, as well as intracellularly produced truncated proteins, are known in the art. For example, such complexes may be produced recombinantly, as described in U.S. Patent No. 6,121,020; Ralston et al., *J. Virol.* (1993) <u>67</u>:6753-6761, Grakoui et al., *J. Virol.* (1993) <u>67</u>:1385-1395; and Lanford et al., *Virology* (1993) <u>197</u>:225-235.

Polynucleotides Encoding the Fusion Proteins and E1E2 Complexes

Polynucleotides contain less than an entire HCV genome and can be RNA or single- or double-stranded DNA. Preferably, the polynucleotides are isolated free of other components, such as proteins and lipids. The polynucleotides encode the fusion proteins, E1 and E2 polypeptides and complexes thereof, described above, and thus comprise coding sequences thereof. Polynucleotides of the invention can also comprise other non-HCV nucleotide sequences, such as sequences coding for linkers, signal sequences, or ligands useful in protein purification such as glutathione-S-transferase and staphylococcal protein A.

Polynucleotides encoding the various HCV polypeptides can be isolated from a genomic library derived from nucleic acid sequences present in, for example, the plasma, serum, or liver homogenate of an HCV infected individual or can be synthesized in the laboratory, for example, using an automatic synthesizer. An

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amplification method such as PCR can be used to amplify polynucleotides from either HCV genomic DNA or cDNA encoding therefor.

Polynucleotides can comprise coding sequences for these polypeptides which occur naturally or can include artificial sequences which do not occur in nature. These polynucleotides can be ligated to form a coding sequence for the fusion proteins and E1E2 complexes using standard molecular biology techniques. If desired, polynucleotides can be cloned into an expression vector and transformed into, for example, bacterial, yeast, insect, or mammalian cells so that the fusion proteins of the invention can be expressed in and isolated from a cell culture.

The expression constructs of the present invention, including the desired fusion, or individual expression constructs comprising the individual components of these fusions, may be used for nucleic acid immunization, to stimulate an immunological response, such as a cellular immune response, using standard gene delivery protocols. Methods for gene delivery are known in the art. See, e.g., U.S. Patent Nos. 5,399,346, 5,580,859, 5,589,466. Genes can be delivered either directly to the vertebrate subject or, alternatively, delivered ex vivo, to cells derived from the subject and the cells reimplanted in the subject. For example, the constructs can be delivered as plasmid DNA, e.g., contained within a plasmid, such as pBR322, pUC, or ColE1

Additionally, the expression constructs can be packaged in liposomes prior to delivery to the cells. Lipid encapsulation is generally accomplished using liposomes which are able to stably bind or entrap and retain nucleic acid. The ratio of condensed DNA to lipid preparation can vary but will generally be around 1:1 (mg DNA:micromoles lipid), or more of lipid. For a review of the use of liposomes as carriers for delivery of nucleic acids, see, Hug and Sleight, *Biochim. Biophys. Acta.* (1991) 1097:1-17; Straubinger et al., in *Methods of Enzymology* (1983), Vol. 101, pp. 512-527.

Liposomal preparations for use with the present invention include cationic (positively charged), anionic (negatively charged) and neutral preparations, with cationic liposomes particularly preferred. Cationic liposomes are readily available. For example, N[1-2,3-dioleyloxy)propyl]-N,N,N-triethylammonium (DOTMA)

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liposomes are available under the trademark Lipofectin, from GIBCO BRL, Grand Island, NY. (See, also, Felgner et al., Proc. Natl. Acad. Sci. USA (1987) 84:7413-7416). Other commercially available lipids include transfectace (DDAB/DOPE) and DOTAP/DOPE (Boerhinger). Other cationic liposomes can be prepared from readily available materials using techniques well known in the art. See, e.g., Szoka et al., Proc. Natl. Acad. Sci. USA (1978) 75:4194-4198; PCT Publication No. WO 90/11092 for a description of the synthesis of DOTAP (1,2-bis(oleoyloxy)-3-(trimethylammonio)propane) liposomes. The various liposome-nucleic acid complexes are prepared using methods known in the art. See, e.g., Straubinger et al., in METHODS OF IMMUNOLOGY (1983), Vol. 101, pp. 512-527; Szoka et al., Proc. Natl. Acad. Sci. USA (1978) 75:4194-4198; Papahadjopoulos et al., Biochim. Biophys. Acta (1975) 394:483; Wilson et al., Cell (1979) 17:77); Deamer and Bangham, Biochim. Biophys. Acta (1976) 443:629; Ostro et al., Biochem. Biophys. Res. Commun. (1977) 76:836; Fraley et al., Proc. Natl. Acad. Sci. USA (1979) 76:3348); Enoch and Strittmatter, Proc. Natl. Acad. Sci. USA (1979) 76:145); Fraley et al., J. Biol. Chem. (1980) 255:10431; Szoka and Papahadiopoulos, Proc. Natl. Acad. Sci. USA (1978) 75:145; and Schaefer-Ridder et al., Science (1982) 215:166.

The DNA can also be delivered in cochleate lipid compositions similar to those described by Papahadjopoulos et al., *Biochem. Biophys. Acta.* (1975) 394:483-491. See, also, U.S. Patent Nos. 4,663,161 and 4,871,488.

A number of viral based systems have been developed for gene transfer into mammalian cells. For example, retroviruses provide a convenient platform for gene delivery systems, such as murine sarcoma virus, mouse mammary tumor virus, Moloney murine leukemia virus, and Rous sarcoma virus. A selected gene can be inserted into a vector and packaged in retroviral particles using techniques known in the art. The recombinant virus can then be isolated and delivered to cells of the subject either *in vivo* or *ex vivo*. A number of retroviral systems have been described (U.S. Patent No. 5,219,740; Miller and Rosman, *BioTechniques* (1989) 7:980-990; Miller, A.D., *Human Gene Therapy* (1990) 1:5-14; Scarpa et al., *Virology* (1991) 180:849-852; Burns et al., *Proc. Natl. Acad. Sci. USA* (1993) 90:8033-8037; and Boris-Lawrie and Temin, *Cur. Opin. Genet. Develop.* (1993) 3:102-109. Briefly,

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retroviral gene delivery vehicles of the present invention may be readily constructed from a wide variety of retroviruses, including for example, B, C, and D type retroviruses as well as spumaviruses and lentiviruses such as FIV, HIV, HIV-1, HIV-2 and SIV (see RNA Tumor Viruses, Second Edition, Cold Spring Harbor

Laboratory, 1985). Such retroviruses may be readily obtained from depositories or collections such as the American Type Culture Collection ("ATCC"; 10801 University Blvd., Manassas, VA 20110-2209), or isolated from known sources using commonly available techniques.

A number of adenovirus vectors have also been described, such as adenovirus Type 2 and Type 5 vectors. Unlike retroviruses which integrate into the host genome, adenoviruses persist extrachromosomally thus minimizing the risks associated with insertional mutagenesis (Haj-Ahmad and Graham, *J. Virol.* (1986) 57:267-274; Bett et al., *J. Virol.* (1993) 67:5911-5921; Mittereder et al., *Human Gene Therapy* (1994) 5:717-729; Seth et al., *J. Virol.* (1994) 68:933-940; Barr et al., *Gene Therapy* (1994) 1:51-58; Berkner, K.L. *BioTechniques* (1988) 6:616-629; and Rich et al., *Human Gene Therapy* (1993) 4:461-476).

Molecular conjugate vectors, such as the adenovirus chimeric vectors described in Michael et al., *J. Biol. Chem.* (1993) <u>268</u>:6866-6869 and Wagner et al., *Proc. Natl. Acad. Sci. USA* (1992) <u>89</u>:6099-6103, can also be used for gene delivery.

Members of the Alphavirus genus, such as but not limited to vectors derived from the Sindbis and Semliki Forest viruses, VEE, will also find use as viral vectors for delivering the gene of interest. For a description of Sindbis-virus derived vectors useful for the practice of the instant methods, see, Dubensky et al., *J. Virol.* (1996) 70:508-519; and International Publication Nos. WO 95/07995 and WO 96/17072.

Other vectors can be used, including but not limited to simian virus 40 and cytomegalovirus. Bacterial vectors, such as Salmonella ssp. Yersinia enterocolitica, Shigella spp., Vibrio cholerae, Mycobacterium strain BCG, and Listeria monocytogenes can be used. Minichromosomes such as MC and MC1, bacteriophages, cosmids (plasmids into which phage lambda cos sites have been inserted) and replicons (genetic elements that are capable of replication under their own control in a cell) can also be used.

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The expression constructs may also be encapsulated, adsorbed to, or associated with, particulate carriers. Such carriers present multiple copies of a selected molecule to the immune system and promote trapping and retention of molecules in local lymph nodes. The particles can be phagocytosed by macrophages and can enhance antigen presentation through cytokine release. Examples of particulate carriers include those derived from polymethyl methacrylate polymers, as well as microparticles derived from poly(lactides) and poly(lactide-co-glycolides), known as PLG. See, e.g., Jeffery et al., *Pharm. Res.* (1993) 10:362-368; and McGee et al., *J. Microencap.* (1996).

One preferred method for adsorbing macromolecules onto prepared microparticles is described in International Publication No. WO 00/050006. Briefly, microparticles are rehydrated and dispersed to an essentially monomeric suspension of microparticles using dialyzable anionic or cationic detergents. Useful detergents include, but are not limited to, any of the various N-methylglucamides (known as MEGAs), such as heptanoyl-N-methylglucamide (MEGA-7), octanoyl-N-methylglucamide (MEGA-8), nonanoyl-N-methylglucamide (MEGA-9), and decanoyl-N-methyl-glucamide (MEGA-10); cholic acid; sodium cholate; deoxycholic acid; sodium deoxycholate; taurocholic acid; sodium taurocholate; taurodeoxycholic acid; sodium taurodeoxycholate; 3-[(3-cholamidopropyl)dimethylammonio] -1-propane-sulfonate (CHAPS); 3-[(3-

cholamidopropyl) dimethylammonio]-2-hydroxy-1-propane-sulfonate (CHAPSO);
-dodecyl-N,N-dimethyl-3-ammonio-1-propane-sulfonate (ZWITTERGENT 3-12);
N,N-bis-(3-D-gluconeamidopropyl)-deoxycholamide (DEOXY-BIGCHAP);
-octylglucoside; sucrose monolaurate; glycocholic acid/sodium glycocholate;
laurosarcosine (sodium salt); glycodeoxycholic acid/sodium glycodeoxycholate;
sodium dodceyl sulfate (SDS); 3-(trimethylsilyl)-1-propanesulfonic acid (DSS);
cetrimide (CTAB, the principal component of which is
hexadecyltrimethylammonium bromide); hexadecyltrimethylammonium bromide;
dodecyltrimethylammonium bromide; hexadecyltrimethyl-ammonium bromide;
tetradecyltrimethylammonium bromide; benzyl dimethyldodecylammonium
bromide; benzyl dimethyl-hexadecylammonium chloride; and benzyl dimethyltetra-

decylammonium bromide. The above detergents are commercially available from e.g., Sigma Chemical Co., St. Louis, MO. Various cationic lipids known in the art can also be used as detergents. See Balasubramaniam et al., 1996, *Gene Ther.*, 3:163-72 and Gao, X., and L. Huang. 1995, *Gene Ther.*, 2:7110-722.

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A wide variety of other methods can be used to deliver the expression constructs to cells. Such methods include DEAE dextran-mediated transfection, calcium phosphate precipitation, polylysine- or polyornithine-mediated transfection, or precipitation using other insoluble inorganic salts, such as strontium phosphate, aluminum silicates including bentonite and kaolin, chromic oxide, magnesium silicate, talc, and the like. Other useful methods of transfection include electroporation, sonoporation, protoplast fusion, liposomes, peptoid delivery, or microinjection. See, e.g., Sambrook et al., *supra*, for a discussion of techniques for transforming cells of interest; and Felgner, P.L., *Advanced Drug Delivery Reviews* (1990) 5:163-187, for a review of delivery systems useful for gene transfer. Methods of delivering DNA using electroporation are described in, e.g., U.S. Patent Nos. 6,132,419; 6,451,002, 6,418,341, 6233,483, U.S. Patent Publication No. 2002/0146831; and International Publication No. WO/0045823.

Moreover, the HCV polynucleotides can be adsorbed to, or entrapped within, an ISCOM. Classic ISCOMs are formed by combination of cholesterol, saponin, phospholipid, and immunogens, such as viral envelope proteins. Generally, the HCV molecules (usually with a hydrophobic region) are solubilized in detergent and added to the reaction mixture, whereby ISCOMs are formed with the HCV molecule incorporated therein. ISCOM matrix compositions are formed identically, but without viral proteins. Proteins with high positive charge may be electrostatically bound in the ISCOM particles, rather than through hydrophobic forces. For a more detailed general discussion of saponins and ISCOMs, and methods of formulating ISCOMs, see Barr et al. (1998) *Adv. Drug Delivery Reviews* 32:247-271 (1998); U.S. Patent Nos. 4,981,684, 5,178,860, 5,679,354 and 6,027,732; European Publ. Nos. EPA 109,942; 180,564 and 231,039; and Coulter et al. (1998) *Vaccine* 16:1243.

Additionally, biolistic delivery systems employing particulate carriers such as gold and tungsten, are especially useful for delivering the expression constructs of

the present invention. The particles are coated with the construct to be delivered and accelerated to high velocity, generally under a reduced atmosphere, using a gun powder discharge from a "gene gun." For a description of such techniques, and apparatuses useful therefore, see, e.g., U.S. Patent Nos. 4,945,050; 5,036,006; 5,100,792; 5,179,022; 5,371,015; and 5,478,744.

Compositions Comprising Fusion Proteins or Polynucleotides

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The invention also provides compositions comprising the fusion proteins or polynucleotides, as well as compositions including the individual components of these fusion proteins or polynucleotides. Compositions of the invention preferably comprise a pharmaceutically acceptable carrier. The carrier should not itself induce the production of antibodies harmful to the host. Pharmaceutically acceptable carriers are well known to those in the art. Such carriers include, but are not limited to, large, slowly metabolized, macromolecules, such as proteins, polysaccharides such as latex functionalized sepharose, agarose, cellulose, cellulose beads and the like, polylactic acids, polyglycolic acids, polymeric amino acids such as polyglutamic acid, polylysine, and the like, amino acid copolymers, and inactive virus particles.

Pharmaceutically acceptable salts can also be used in compositions of the invention, for example, mineral salts such as hydrochlorides, hydrobromides, phosphates, or sulfates, as well as salts of organic acids such as acetates, proprionates, malonates, or benzoates. Especially useful protein substrates are serum albumins, keyhole limpet hemocyanin, immunoglobulin molecules, thyroglobulin, ovalbumin, tetanus toxoid, and other proteins well known to those of skill in the art. Compositions of the invention can also contain liquids or excipients, such as water, saline, glycerol, dextrose, ethanol, or the like, singly or in combination, as well as substances such as wetting agents, emulsifying agents, or pH buffering agents. Liposomes can also be used as a carrier for a composition of the invention, such liposomes are described above.

If desired, co-stimulatory molecules which improve immunogen presentation to lymphocytes, such as B7-1 or B7-2, or cytokines such as GM-CSF, IL-2, and IL-

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12, can be included in a composition of the invention. Optionally, adjuvants can also be included in a composition. Adjuvants which can be used include, but are not limited to: (1) aluminum salts (alum), such as aluminum hydroxide, aluminum phosphate, aluminum sulfate, etc.; (2) oil-in-water emulsion formulations (with or without other specific immunostimulating agents such as muramyl peptides (see below) or bacterial cell wall components), such as for example (a) MF59 (U.S. Patent No. 6.299,884; Chapter 10 in Vaccine design: the subunit and adjuvant approach, eds. Powell & Newman, Plenum Press 1995), containing 5% Squalene, 0.5% TWEEN 80TM, and 0.5% SPAN 85TM (optionally containing various amounts of MTP-PE (see below), although not required) formulated into submicron particles using a microfluidizer such as Model 110Y microfluidizer (Microfluidics, Newton, MA), (b) SAF, containing 10% Squalane, 0.4% TWEEN 80TM, 5% pluronic-blocked polymer L121, and thr-MDP either microfluidized into a submicron emulsion or vortexed to generate a larger particle size emulsion, and (c) RIBITM adjuvant system (RAS), (Ribi Immunochem, Hamilton, MT) containing 2% Squalene, 0.2% TWEEN 80TM, and one or more bacterial cell wall components from the group consisting of monophosphorylipid A (MPL), trehalose dimycolate (TDM), and cell wall skeleton (CWS), preferably MPL + CWS (DETOXTM); (3) saponin adjuvants, such as QS21 or STIMULONTM (Cambridge Bioscience, Worcester, MA) may be used or particles generated therefrom such as ISCOMs (immunostimulating complexes), which ISCOMs may be devoid of additional detergent, see, e.g., International Publication No. WO 00/07621; (4) Complete Freund's Adjuvant (CFA) and Incomplete Freund's Adjuvant (IFA); (5) cytokines, such as interleukins (IL-1, IL-2, IL-4, IL-5, IL-6, IL-7, IL-12 (International Publication No. WO 99/44636), etc.), interferons (e.g., gamma interferon), macrophage colony stimulating factor (M-CSF), tumor necrosis factor (TNF), etc.; (6) detoxified mutants of a bacterial ADP-ribosylating toxin such as a cholera toxin (CT), a pertussis toxin (PT), or an E. coli heat-labile toxin (LT), particularly LT-K63 (where lysine is substituted for the wild-type amino acid at position 63) LT-R72 (where arginine is substituted for the wild-type amino acid at position 72), CT-S109 (where serine is substituted for the wild-type amino acid at position 109), and PT-K9/G129 (where lysine is substituted for the wild-type amino

acid at position 9 and glycine substituted at position 129) (see, e.g., International Publication Nos. W093/13202 and W092/19265); (7) MPL or 3-O-deacylated MPL (3dMPL) (see, e.g., GB 2220221), EP-A-0689454, optionally in the substantial absence of alum when used with pneumococcal saccharides (see, e.g., International Publication No. WO 00/56358); (8) combinations of 3dMPL with, for example, 5 OS21 and/or oil-in-water emulsions (see, e.g., EP-A-0835318, EP-A-0735898, EP-A-0761231; (9) oligonucleotides comprising CpG motifs (see, e.g., Roman et al. (1997) Nat. Med. 3:849-854; Weiner et al. (1997) Proc. Natl. Acad. Sci. USA 94:10833-10837; Davis et al. (1998) J. Immunol. 160:870-876; Chu et al. (1997) J. Exp. Med. 186:1623-1631; Lipford et al. (1997) Eur. J. Immunol. 27:2340-2344; 10 Moldoveanu et al. (1988) Vaccine 16:1216-1224; Krieg et al. (1995) Nature 374:546-549; Klinman et al. (1996) Proc. Natl. Acad. Sci. USA 93:2879-2883; Ballas et al. (1996) J. Immunol. 157:1840-1845; Cowdery et al. (1996) J. Immunol. 156:4570-4575; Halpern et al. (1996) Cell Immunol. 167:72-78; Yamamoto et al. (1988) Jpn. J. Cancer Res. 79:866-873; Stacey et al. (1996) J. Immunol. 157:2116-15 2122; Messina et al. (1991) J. Immunol. 147:1759-1764; Yi et al. (1996) J. Immunol. 157:4918-4925; Yi et al. (1996) J. Immunol. 157:5394-5402; Yi et al. (1998) J. Immunol. 160:4755-4761; Yi et al. (1998) J. Immunol. 160:5898-5906; International Publication Nos. WO 96/02555, WO 98/16247, WO 98/18810, WO 98/40100, WO 98/55495, WO 98/37919 and WO 98/52581), such as those containing at least on CG 20 dinucleotide, with cytosine optionally replaced with 5-methylcytosine; (10) a polyoxyethylene ether or a polyoxyethylene ester (see, e.g., International Publication No. WO 99/52549); (11) a polyoxyethylene sorbitan ester surfactant in combination with an octoxynol (see, e.g., International Publication No. WO 01/21207) or a polyoxyethylene alkyl ether or ester surfactant in combination with at least one 25 additional non-ionic surfactant such as an octoxynol (see, e.g., International Publication No. WO 01/21152); (12) a saponin and an immunostimulatory oligonucleotide such as a CpG oligonucleotide (see, e.g., International Publication No. WO 00/62800); (13) an immunostimulant and a particle of metal salt (see, e.g., International Publication No. WO 00/23105); and (14) other substances that act as 30 immunostimulating agents to enhance the effectiveness of the composition.

Muramyl peptides include, but are not limited to, N-acetyl-muramyl-L-threonyl-D-isoglutamine (thr-MDP), N-acteyl-normuramyl-L-alanyl-D-isoglutame (nor-MDP), -acetylmuramyl-L-alanyl-D-isoglutaminyl-L-alanine-2-(l'-2'-dipalmitoyl-sn-glycero-3-huydroxyphosphoryloxy)-ethylamine (MTP-PE), etc.

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Moreover, the HCV proteins can be adsorbed to, or entrapped within, an ISCOM, as described above. Additionally, ISCOMs with adsorbed HCV core proteins, either the entire core region or a fragment of HCV core protein, may be added to the formulations. Most preferably, the HCV core protein is a fragment comprising a polypeptide from the region spanning amino acid positions 121-135. See, e.g., International Publication No. WO 01/37869A.

As explained above, the composition may also contain immunostimulatory molecules, either in addition to or in place of the antigen delivery system.

Immunostimulatory agents for use herein include, without limitation, monophosphorylipid A (MPL), trehalose dimycolate (TDM), and cell wall skeleton (CWS), preferably MPL + CWS (DetoxTM). MPL may be formulated into an emulsion to enhance its immunostimulatory affect. See, e.g., Ulrich et al., "MPLr immunostimulat: adjuvant formulations." in Vaccine Adjuvants: Prepartion Methods and Research Protocols (O'Hagan DT, ed.) Human Press Inc., NJ (2000) pp. 273-282. MPL has been shown to induce the synthesis and release of cytokines, particularly IL-2 and IFN-γ. Other useful immunostimulatory molecules include LPS and immunostimulatory nucleic acid sequences (ISS), including but not limited to, unmethylated CpG motifs, such as CpG oligonucleotides.

Oligonucleotides containing unmethylated CpG motifs have been shown to induce activation of B cells, NK cells and antigen-presenting cells (APCs), such as monocytes and macrophages. See, e.g., U.S. Patent No. 6,207,646. Thus, adjuvants derived from the CpG family of molecules, CpG dinucleotides and synthetic oligonucleotides which comprise CpG motifs (see, e.g., Krieg et al. *Nature* (1995) 374:546 and Davis et al. *J. Immunol.* (1998) 160:870-876) such as any of the various immunostimulatory CpG oligonucleotides disclosed in U.S. Patent No. 6,207,646, may be used in the subject methods and compositions. Such CpG oligonucleotides generally comprise at least 8 up to about 100 basepairs, preferably 8 to 40 basepairs,

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more preferably 15-35 basepairs, preferably 15-25 basepairs, and any number of basepairs between these values. For example, oligonucleotides comprising the consensus CpG motif, represented by the formula 5'-X₁CGX₂-3', where X₁ and X₂ are nucleotides and C is unmethylated, will find use as immunostimulatory CpG molecules. Generally, X₁ is A, G or T, and X₂ is C or T. Other useful CpG molecules include those captured by the formula $5'-X_1X_2CGX_3X_4$, where X_1 and X_2 are a sequence such as GpT, GpG, GpA, ApA, ApT, ApG, CpT, CpA, CpG, TpA, TpT or TpG, and X₃ and X₄ are TpT, CpT, ApT, ApG, CpG, TpC, ApC, CpC, TpA, ApA, GpT, CpA, or TpG, wherein "p" signifies a phosphate bond. Preferably, the oligonucleotides do not include a GCG sequence at or near the 5'- and/or 3' terminus. Additionally, the CpG is preferably flanked on its 5'-end with two purines (preferably a GpA dinucleotide) or with a purine and a pyrimidine (preferably, GpT), and flanked on its 3'-end with two pyrimidines, preferably a TpT or TpC dinucleotide. Thus, preferred molecules will comprise the sequence GACGTT, GACGTC, GTCGTT or GTCGCT, and these sequences will be flanked by several additional nucleotides. The nucleotides outside of this central core area appear to be extremely amendable to change.

Moreover, the CpG oligonucleotides for use herein may be double- or single-stranded. Double-stranded molecules are more stable *in vivo* while single-stranded molecules display enhanced immune activity. Additionally, the phosphate backbone may be modified, such as phosphorodithioate-modified, in order to enhance the immunostimulatory activity of the CpG molecule. As described in U.S. Patent No. 6,207,646, CpG molecules with phosphorothioate backbones preferentially activate B-cells, while those having phosphodiester backbones preferentially activate monocytic (macrophages, dendritic cells and monocytes) and NK cells.

One exemplary CpG oligonucleotide for use in the present compositions has the sequence 5'-TCCATGACGTTCCTGACGTT-3' (SEQ ID NO:6).

CpG molecules can readily be tested for their ability to stimulate an immune response using standard techniques, well known in the art. For example, the ability of the molecule to stimulate a humoral and/or cellular immune response is readily determined using the immunoassays described above. Moreover, the antigen and

adjuvant compositions can be administered with and without the CpG molecule to determine whether an immune response is enhanced.

The HCV proteins may also be encapsulated, adsorbed to, or associated with, particulate carriers, as described above with reference to the HCV polynucleotides.

5 As explained above, examples of particulate carriers include those derived from polymethyl methacrylate polymers, as well as microparticles derived from poly(lactides) and poly(lactide-co-glycolides), known as PLG. See, e.g., Jeffery et al., *Pharm. Res.* (1993) 10:362-368; and McGee et al., *J. Microencap.* (1996). One preferred method for adsorbing macromolecules onto prepared microparticles is described above and in International Publication No. WO 00/050006.

Methods of Producing HCV-Specific Antibodies

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The HCV proteins can be used to produce HCV-specific polyclonal and monoclonal antibodies. HCV-specific polyclonal and monoclonal antibodies specifically bind to HCV antigens. Polyclonal antibodies can be produced by administering the fusion protein to a mammal, such as a mouse, a rabbit, a goat, or a horse. Serum from the immunized animal is collected and the antibodies are purified from the plasma by, for example, precipitation with ammonium sulfate, followed by chromatography, preferably affinity chromatography. Techniques for producing and processing polyclonal antisera are known in the art.

Monoclonal antibodies directed against HCV-specific epitopes present in the proteins can also be readily produced. Normal B cells from a mammal, such as a mouse, immunized with an HCV protein, can be fused with, for example, HAT-sensitive mouse myeloma cells to produce hybridomas. Hybridomas producing HCV-specific antibodies can be identified using RIA or ELISA and isolated by cloning in semi-solid agar or by limiting dilution. Clones producing HCV-specific antibodies are isolated by another round of screening.

Antibodies, either monoclonal and polyclonal, which are directed against HCV epitopes, are particularly useful for detecting the presence of HCV or HCV antigens in a sample, such as a serum sample from an HCV-infected human. An immunoassay for an HCV antigen may utilize one antibody or several antibodies. An

immunoassay for an HCV antigen may use, for example, a monoclonal antibody directed towards an HCV epitope, a combination of monoclonal antibodies directed towards epitopes of one HCV polypeptide, monoclonal antibodies directed towards epitopes of different HCV polypeptides, polyclonal antibodies directed towards the same HCV antigen, polyclonal antibodies directed towards different HCV antigens, or a combination of monoclonal and polyclonal antibodies. Immunoassay protocols may be based, for example, upon competition, direct reaction, or sandwich type assays using, for example, labeled antibody. The labels may be, for example, fluorescent, chemiluminescent, or radioactive.

The polyclonal or monoclonal antibodies may further be used to isolate HCV particles or antigens by immunoaffinity columns. The antibodies can be affixed to a solid support by, for example, adsorption or by covalent linkage so that the antibodies retain their immunoselective activity. Optionally, spacer groups may be included so that the antigen binding site of the antibody remains accessible. The immobilized antibodies can then be used to bind HCV particles or antigens from a biological sample, such as blood or plasma. The bound HCV particles or antigens are recovered from the column matrix by, for example, a change in pH.

HCV-Specific T cells

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HCV-specific T cells that are activated by the above-described fusions and E1E2 complexes, including the NS3NS4NS5a fusion protein or NS3NS4NS5aNS5b fusion protein, and the E1E2 complexes, expressed *in vivo* or *in vitro*, preferably recognize an epitope of an HCV polypeptide such as an E1, E2, NS3, NS4, NS5a, NS5b polypeptide, including an epitope of an NS3NS4NS5a fusion protein or an NS3NS4NS5aNS5b fusion protein, or an E1E2 complex. HCV-specific T cells can be CD8+ or CD4+.

HCV-specific CD8⁺ T cells preferably are cytotoxic T lymphocytes (CTL) which can kill HCV-infected cells that display E1, E2, NS3, NS4, NS5a, NS5b epitopes complexed with an MHC class I molecule. HCV-specific CD8⁺ T cells may also express interferon-γ (IFN-γ). HCV-specific CD8⁺ T cells can be detected by, for example, ⁵¹Cr release assays (see the examples). ⁵¹Cr release assays measure the

ability of HCV-specific CD8⁺ T cells to lyse target cells displaying an E1, E2, E1E2, NS3, NS4, NS5a, NS5b, NS3NS4NS5a, or NS3NS4NS5aNS5b epitope. HCV-specific CD8⁺ T cells which express IFN-γ can also be detected by immunological methods, preferably by intracellular staining for IFN-γ after *in vitro* stimulation with an E1, E2, NS3, an NS4, an NS5a, or an NS5b polypeptide (see the examples).

HCV-specific CD4⁺ cells activated by the above-described E1E2 complexes and fusions, such as an E1 polypeptide, an E2 polypeptide, an E1E2 complex, NS3NS4NS5a or NS3NS4NS5aNS5b fusion protein, expressed *in vivo* or *in vitro*, preferably recognize an epitope of an E1, E2, NS3, NS4, NS5a, or NS5b polypeptide, including an epitope of an E1E2 complex, NS3NS4NS5a or NS3NS4NS5aNS5b fusion protein, that is bound to an MHC class II molecule on an HCV-infected cell and proliferate in response to stimulating E1E2 complexes with NS3NS4NS5a or NS3NS4NS5a or NS3NS4NS5aNS5b peptides, with or without a core polypeptide.

HCV-specific CD4⁺ T cells can be detected by a lymphoproliferation assay (see examples). Lymphoproliferation assays measure the ability of HCV-specific CD4⁺ T cells to proliferate in response to an E1, E2, NS3, an NS4, an NS5a, or an NS5b epitope.

Methods of Activating HCV-Specific T Cells.

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The HCV proteins or polynucleotides can be used to stimulate an immune response, such as to activate HCV-specific T cells either *in vitro* or *in vivo*.

Activation of HCV-specific T cells can be used, *inter alia*, to provide model systems to optimize CTL responses to HCV and to provide prophylactic or therapeutic treatment against HCV infection. For *in vitro* activation, proteins are preferably supplied to T cells via a plasmid or a viral vector, such as an adenovirus vector, as described above.

Polyclonal populations of T cells can be derived from the blood, and preferably

from peripheral lymphoid organs, such as lymph nodes, spleen, or thymus, of mammals that have been infected with an HCV. Preferred mammals include mice, chimpanzees, baboons, and humans. The HCV serves to expand the number of

activated HCV-specific T cells in the mammal. The HCV-specific T cells derived from the mammal can then be restimulated *in vitro* by adding, e.g., HCV E1E2 and NS3NS4NS5a or NS3NS4NS5aNS5b epitopic peptides, with or without a core polypeptide, to the T cells. The HCV-specific T cells can then be tested for, *inter alia*, proliferation, the production of IFN-γ, and the ability to lyse target cells displaying E1E2, NS3NS4NS5a or NS3NS4NS5aNS5b epitopes *in vitro*.

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In a lymphoproliferation assay (see examples), HCV-activated CD4⁺ T cells proliferate when cultured with an NS3, NS4, NS5a, NS5b, NS3NS4NS5a, or NS3NS4NS5aNS5b epitopic peptide, but not in the absence of an epitopic peptide.

Thus, particular E1, E2, NS3, NS4, NS5a, NS5b, NS3NS4NS5a and NS3NS4NS5aNS5b epitopes that are recognized by HCV-specific CD4⁺ T cells can be identified using a lymphoproliferation assay.

Similarly, detection of IFN-γ in HCV-specific CD8⁺ T cells after *in vitro* stimulation with the above-described HCV proteins, can be used to identify E1, E2, E1E2, NS3, NS4, NS5a, NS5b, NS3NS4NS5a, and NS3NS4NS5aNS5b epitopes that particularly effective at stimulating CD8⁺ T cells to produce IFN-γ (see examples).

Further, ⁵¹Cr release assays are useful for determining the level of CTL response to HCV. See Cooper et al. Immunity 10:439-449. For example, HCV-specific CD8⁺ T cells can be derived from the liver of an HCV infected mammal. These T cells can be tested in ⁵¹Cr release assays against target cells displaying, e.g., E1E2, NS3NS4NS5a and/or NS3NS4NS5aNS5b epitopes. Several target cell populations expressing different E1E2, NS3NS4NS5a and/or NS3NS4NS5aNS5b epitopes can be constructed so that each target cell population displays different epitopes of E1E2, NS3NS4NS5a and/or NS3NS4NS5aNS5b. The HCV-specific CD8⁺ cells can be assayed against each of these target cell populations. The results of the ⁵¹Cr release assays can be used to determine which epitopes of E1E2, NS3NS4NS5a and/or NS3NS4NS5aNS5b are responsible for the strongest CTL response to HCV. E1E2 complexes, NS3NS4NS5a fusion proteins or

NS3NS4NS5aNS5b fusion proteins, with or without core polypeptides, which

contain the epitopes responsible for the strongest CTL response can then be constructed using the information derived from the ⁵¹Cr release assays.

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HCV proteins as described above, or polynucleotides encoding such proteins, can be administered to a mammal, such as a mouse, baboon, chimpanzee, or human, to stimulate an immune response, such as to activate HCV-specific T cells *in vivo*. Administration can be by any means known in the art, including parenteral, intranasal, intramuscular or subcutaneous injection, including injection using a biological ballistic gun ("gene gun"), as discussed above.

Preferably, injection of an HCV polynucleotide is used to activate T cells. In addition to the practical advantages of simplicity of construction and modification, injection of the polynucleotides results in the synthesis of a fusion protein in the host. Thus, these immunogens are presented to the host immune system with native post-translational modifications, structure, and conformation. The polynucleotides are preferably injected intramuscularly to a large mammal, such as a human, at a dose of 0.5, 0.75, 1.0, 1.5, 2.0, 2.5, 5 or 10 mg/kg.

A composition of the invention comprising the HCV proteins or polynucleotides is administered in a manner compatible with the particular composition used and in an amount which is effective to stimulate an immune response, such as to activate HCV-specific T cells as measured by, *inter alia*, a ⁵¹Cr release assay, a lymphoproliferation assay, or by intracellular staining for IFN-γ. The proteins and/or polynucleotides can be administered either to a mammal which is not infected with an HCV or can be administered to an HCV-infected mammal. The particular dosages of the polynucleotides or proteins in a composition will depend on many factors including, but not limited to the species, age, and general condition of the mammal to which the composition is administered, and the mode of administration of the composition. An effective amount of the composition of the invention can be readily determined using only routine experimentation. *In vitro* and *in vivo* models described above can be employed to identify appropriate doses. The amount of polynucleotide used in the example described below provides general guidance which can be used to optimize the activation of HCV-specific T cells either

in vivo or in vitro. Generally, 0.5, 0.75, 1.0, 1.5, 2.0, 2.5, 5 or 10 mg of an HCV fusion and E1 and E2 polypeptides, such as an E1E2 complex, an NS3NS4NS5a or NS3NS4NS5b fusion protein or polynucleotide, with or without a core polypeptide, will be administered to a large mammal, such as a baboon, chimpanzee, or human. If desired, co-stimulatory molecules or adjuvants can also be provided before, after, or together with the compositions.

Immune responses of the mammal generated by the delivery of a composition of the invention, including activation of HCV-specific T cells, can be enhanced by varying the dosage, route of administration, or boosting regimens. Compositions of the invention may be given in a single dose schedule, or preferably in a multiple dose schedule in which a primary course of vaccination includes 1-10 separate doses, followed by other doses given at subsequent time intervals required to maintain and/or reenforce an immune response, for example, at 1-4 months for a second dose, and if needed, a subsequent dose or doses after several months.

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Deposits of Strains Useful in Practicing the Invention

A deposit of biologically pure cultures of the following strains was made with the American Type Culture Collection, 10801 University Boulevard, Manassas, VA. The accession number indicated was assigned after successful viability testing, and the requisite fees were paid. made under the provisions of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purpose of Patent Procedure and the Regulations thereunder (Budapest Treaty). This assures maintenance of viable cultures for a period of thirty (30) years from the date of deposit. The organisms will be made available by the ATCC under the terms of the Budapest Treaty, which assures permanent and unrestricted availability of the progeny to one determined by the U.S. Commissioner of Patents and Trademarks to be entitled thereto according to 35 U.S.C. §122 and the Commissioner's rules pursuant thereto (including 37 C.F.R. §1.12 with particular reference to 886 OG 638). Upon the granting of a patent, all restrictions on the availability to the public of the deposited cultures will be irrevocably removed.

These deposits are provided merely as convenience to those of skill in the art, and are not an admission that a deposit is required under 35 U.S.C. §112. The nucleic acid sequences of these genes, as well as the amino acid sequences of the molecules encoded thereby, are incorporated herein by reference and are controlling in the event of any conflict with the description herein. A license may be required to make, use, or sell the deposited materials, and no such license is hereby granted.

	Plasmid	Deposit Date	ATCC No.
10	E1E2-809	August 16, 2001	PTA-3643

III. Experimental

Below are examples of specific embodiments for carrying out the present invention. The examples are offered for illustrative purposes only, and are not intended to limit the scope of the present invention in any way. Those of skill in the art will readily appreciate that the invention may be practiced in a variety of ways given the teaching of this disclosure.

Efforts have been made to ensure accuracy with respect to numbers used (e.g., amounts, temperatures, etc.), but some experimental error and deviation should, of course, be allowed for.

EXAMPLE 1

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Production of NS3NS4NS5a Polynucleotides.

A polynucleotide encoding NS3NS4NS5a (approximately amino acids 1027 to 2399, numbered relative to HCV-1) (also termed "NS345a" herein) or NS5a (approximately amino acids 1973 to 2399, numbered relative to HCV-1) was isolated from an HCV. Polynucleotides encoding a methionine residue were ligated to the 5' end of these polynucleotides and the polynucleotides were cloned into plasmid, vaccinia virus, and adenovirus vectors.

Immunization Protocols. In one immunization protocol, mice were immunized with 50 μ g of plasmid DNA encoding either NS5a or encoding an NS3NS4NS5a fusion protein by intramuscular injection into the tibialis anterior. A booster injection of 10^7 pfu of vaccinia virus (VV)-NS5a (intraperitoneal) or 50 μ g of plasmid control (intramuscular) was provided 6 weeks later.

In another immunization protocol, mice were injected intramuscularly in the tibialis anterior with 10¹⁰ adenovirus particles encoding an NS3NS4NS5a fusion protein. An intraperitoneal booster injection of 10⁷ pfu of VV-NS5a or an intramuscular booster injection of 10¹⁰ adenovirus particles encoding NS3NS4NS5a was provided 6 weeks later.

EXAMPLE 2

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Immunization with DNA encoding an NS3NS4NS5a fusion protein activates HCV-specific CD8⁺ T cells.

HCV-specific T cells to lyse target cells displaying an NS5a epitope. Spleen cells were pooled from the immunized animals. These cells were restimulated *in vitro* for 6 days with the CTL epitopic peptide p214K9 (2152-HEYPVGSQL-2160; SEQ ID NO:1) from HCV-NS5a in the presence of IL-2. The spleen cells were then assayed for cytotoxic activity in a standard ⁵¹Cr release assay against peptide-sensitized target cells (L929) expressing class I, but not class II MHC molecules, as described in Weiss (1980) J. Biol. Chem. 255:9912-9917. Ratios of effector (T cells) to target (B cells) of 60:1, 20:1, and 7:1 were tested. Percent specific lysis was calculated for each effector to target ratio.

The results of the assays are shown in Tables 1 and 2. Table 1 demonstrates that immunization with plasmid DNA encoding an NS3NS4NS5a fusion protein activates CD8⁺ T cells which recognize and lyse target cells displaying an NS5a epitope. Surprisingly the NS5a polypeptide of the NS3NS4NS5a fusion protein was able to activate T cells even though the NS5a polypeptide was present in a fusion protein.

Similarly, Table 2 demonstrates that delivery of the NS3NS4NS5a fusion protein to mice by means of an adenovirus vector also activates CD8⁺ T cells which recognize and lyse target cells displaying an HCV NS5a epitope. Thus, immunization with either "naked" (plasmid) DNA encoding an NS3NS4NS5a fusion protein or adenovirus vector-encoded fusion protein can be used to activate HCV-specific T cells.

EXAMPLE 3

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Immunization with DNA encoding an NS3NS4NS5a fusion protein activates HCV-specific CD8⁺ T cells which express IFN-γ.

Intracellular Staining for Interferon-gamma (IFN-γ). Intracellular staining for IFN-γ was used to identify the CD8⁺ T cells that secrete IFN-γ after *in vitro* stimulation with the NS5a epitope p214K9. Spleen cells of individual immunized mice were restimulated *in vitro* either with p214K9 or with a non-specific peptide for 6-12 hours in the presence of IL-2 and monensin. The cells were then stained for surface CD8 and for intracellular IFN-γ and analyzed by flow cytometry. The percent of CD8⁺ T cells which were also positive for IFN-γ was then calculated. The results of these assays are shown in Tables 1 and 2. Table 1 demonstrates that CD8⁺ T cells activated in response to immunization with plasmid DNA encoding an NS3NS4NS5a fusion protein also express IFN-γ. Immunization with an NS3NS4NS5a fusion protein encoded in an adenovirus also results in CD8⁺ HCV-specific T cells which express IFN-γ, although to a lesser extent than immunization with a plasmid-encoded NS3NS4NS5a fusion protein (Table 2).

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	able 1. HCV	/-NS5a-Sp	ecific CD8+	Table 1. HCV-NS5a-Specific CD8+ T Cells in Mice Immunized with NS5a or NS345a DNA	e Immunized	with NS5a o	r NS345a D	NA
	D ₁₈	51Cr Release Assay	Assay		n	Intracellular Staining for IFN-y	aining for IF	N-y
	Percent Sp	ecific Lysi	Percent Specific Lysis of Targets		Percent o	Percent of CD8+ T Cells Positive for IFN-g**	Ils Positive	for IFN-g
E:T ratio	NS5a DNA	NA	NS34	NS345a DNA	NS5a DNA	DNA	NS34	NS345a DNA
	p214K9	1	p214K9	-	p214K9	•	p214K9	•
60:1	77	5	99	9				·
20:1	61	4	49	2	1.74	0.26	1.18	0.40
7:1	29	1	29	4				

*Target cells (L929) were pulsed with p214K9 or media alone and labeled with ⁵¹Cr. *Spleen cells were cultured with p214K9 or media alone for 12 hours in the presence of monensin.

p214K9 is a CTL epitopic peptide (2152-HEYPVGSQL-2160, SEQ ID NO:1) from HCV-NS5a '-' refers to the absence of peptide

							
S345a	N-y	for IFN-g	NS345a DNA	p214J		0.00	
coding for N	aining for IF	ells Positive	NS34	p214K9		0.25	
or DNA En	Intracellular Staining for IFN-y	Percent of CD8+ T Cells Positive for IFN-g**	Adeno	p214J		0.13	
y Adenovirus	In	Percent o	NS345a Adeno	p214K9		3.24	
Table 2. HCV-NS5a-Specific CD8+ T Cells Primed by Adenovirus or DNA Encoding for NS345a			Adeno NS345a DNA	•	5	3	3
ific CD8+ T	Release Assay	Percent Specific Lysis of Targets*		-p214K9	55	22	10
S5a-Specific				Adeno	•	2	, 7
le 2. HCV-h	51C		NS345a Adeno	p214K9	92	85	62
Tab			E:T ratio		60:1	20:1	7:1

*Target cells (L929) were pulsed with p214K9 or p214J and labeled with ⁵¹Cr. *Spleen cells were cultured with p214K9 or p214J for 12 hours in the presence of monensin.

p214K9 is a CTL epitopic peptide (2152-HEYPVGSQL-2160, SEQ ID NO:1) from HCV-NS5a P214J is a control peptide (10 mer) from HCV-NS5a

EXAMPLE 4

Immunization with DNA encoding an NS3NS4NS5a fusion protein stimulates proliferation of HCV-specific CD4⁺ T cells.

Lymphoproliferation assay. Spleen cells from pooled immunized mice were depleted of CD8⁺ T cells using magnetic beads and were cultured in triplicate with either p222D, an NS5a-epitopic peptide from HCV-NS5a (2224-AELIEANLLWRQEMG-2238; SEQ ID NO:2), or in medium alone. After 72 hours, cells were pulsed with 1μ Ci per well of ³H-thymidine and harvested 6-8 hours later. Incorporation of radioactivity was measured after harvesting. The mean cpm was calculated.

As shown in Table 3, immunization with a plasmid-encoded NS3NS4NS5a fusion protein stimulates proliferation of CD4⁺ HCV-specific T cells. Immunization with an adenovirus vector encoding the fusion protein also resulted in stimulated proliferation of CD4⁺ HCV-specific T cells (Table 4).

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		ecific CD4+ T C NS5a or NS345a			
	Mea	an CPM			
NS5a	DNA	NS34	5a DNA		
p222D	media	p222D media			
4523	740	4562	861		
(x6	5.1)	()	(5.3)		

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p222D is a CD4+ epitopic peptide (aa: 2224-AELIEANLLWRQEMG-2238, SEQ ID NO:2) from HCV-NS5a

		cific CD4+ T Cel A Encoding for N	
	Me	an CPM	
NS345a	Adeno	NS34	5a DNA
p222D	media	p222D	media
896	357	1510	385
(x2	2.5)	(х	:3.9)

p222D is a CD4+ epitopic peptide (aa: 2224-AELIEANLLWRQEMG-2238, SEQ ID NO:2) from HCV-NS5a

EXAMPLE 5

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Efficiency of NS345a-encoding DNA Vaccine Formulations to prime CTLs in mice.

Mice were immunized with either 10-100 μg of plasmid DNA encoding NS345a fusion protein as described in Example 1, with PLG-linked DNA encoding NS345a, described below, or with DNA encoding NS345a, delivered via electroporation (see, e.g., U.S. Patent Nos. 6,132,419; 6,451,002, 6,418,341, 6233,483, U.S. Patent Publication No. 2002/0146831; and International Publication No. WO/0045823, for this delivery technique). The immunizations were followed by a booster injection 6 weeks later of 1 x 10⁷ pfu vaccinia virus encoding NS5a, plasmid DNA encoding NS345a or plasmid DNA encoding NS5a each as described in Example 1.

PLG-delivered DNA. The polylactide-co-glycolide (PLG) polymers were obtained from Boehringer Ingelheim, U.S.A. The PLG polymer used in this study was RG505, which has a copolymer ratio of 50/50 and a molecular weight of 65 kDa (manufacturers data). Cationic microparticles with adsorbed DNA were prepared using a modified solvent evaporation process, essentially as described in Singh et al., Proc. Natl. Acad. Sci. USA (2000) 97:811-816. Briefly, the microparticles were prepared by emulsifying 10 ml of a 5% w/v polymer solution in methylene chloride with 1 ml of PBS at high speed using an IKA homogenizer. The primary emulsion was then added to 50ml of distilled water containing cetyl trimethyl ammonium

bromide (CTAB) (0.5% w/v). This resulted in the formation of a w/o/w emulsion which was stirred at 6000 rpm for 12 hours at room temperature, allowing the methylene chloride to evaporate. The resulting microparticles were washed twice in distilled water by centrifugation at 10,000 g and freeze dried. Following preparation, washing and collection, DNA was adsorbed onto the microparticles by incubating 100 mg of cationic microparticles in a 1mg/ml solution of DNA at 4 C for 6 hours. The microparticles were then separated by centrifugation, the pellet washed with TE buffer and the microparticles were freeze dried.

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CTL activity and IFN-γ expression were measured by ⁵¹Cr release assay or intracellular staining as described in examples 2 and 3 respectively. The results are shown in Table 5.

Results demonstrate that immunization using plasmid DNA encoding for NS345a to prime mice results in activation of CD8+ HCV specific T cells.

	Table 5: E	fficiency of	NS345a-I		NA Vacc	ine Formu	lations	to Prime C	TLs in
					or IFN-gar cells that a g+)				
5	NS345a DNA Vaccines	Boost	Mean	SdtdevP	# of mice tested	% respon d- ing	# of expts	fold increase vs. 'naked' DNA	CTL activ i- ty?
	NS345a DNA	VVNS5a	1.02	1.70	41	68%	10	N/A	YES
10	NS345a DNA	NS345a DNA	0.02	0.04	22	5%	5	N/A	YES
	NS345a DNA	NS5a DNA	0.22	0.21	24	63%	5	N/A	YES
15	NS345a DNA eV (electro- poration)	VVNS5a	5.00	4.36	7	100%	2	4.90	YES
	PLGNS34 5a DNA	VVNS5a	2.65	2.54	6	100%	2	2.60	YES
20	PLGNS34 5a DNA	NS5a DNA	0.33	0.24	15	80%	3	1.50	YES

EXAMPLE 6

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Immunization routes and replicon particles SINCR (DC+) encoding for NS345a

Alphavirus replicon particles, for example, SINCR (DC+) were prepared as

described in Polo et al., Proc. Natl. Acad. Sci. USA (1999) 96:4598-4603. Mice were
injected with 5 x 10⁶ IU SINCR (DC+) replicon particles encoding for NS345a
intramuscularly (IM) as described in Example 1, or subcutaneously (S/C) at the base of the
tail (BoT) and foot pad (FP), or with a combination of 2/3 of the DNA delivered via IM
administration and 1/3 via a BoT route. The immunizations were followed by a booster
injection of vaccinia virus encoding NS5a as described in Example 1.

IFN-γ expression was measured by intracellular staining as described in Example 3. The results are shown in Table 6. The results demonstrate that immunization via SINCR (DC+) replicon particles encoding for NS345a by a variety of routes results in CD8+ HCV specific T cells which express IFN-γ.

able 6:	Table 6: Immunization Routes and SINCR (DC+) Replicon Particles Encoding NS345a (all mice VVNS5a challenged)	es and SINCR (DC+) Replicon (all mice VVNS5a challenged)	Replicon Partic Ilenged)	les Encoding	NS345a	
			ICS for IFN-gamma (%CD8+ cells that are IFN-g+)	N-gamma hat are IFN-g	(+	
Immunization Route	te	Mean	SdtdevP	# of mice tested	# of expts	% responding mice
100% IM (ta)		1.11	0.63	3	1	100%
100% S/C (BoT + FP)	FP)	0.62	0.29	3	1	.100%
2/3 IM (ta) + 1/3 S/C (BoT)	(BoT)	2.43	2.00	3		100%

EXAMPLE 7

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SINCR (DC+) vs SINDC (LP) replicon particles encoding for NS345a

Alphavirus replicon particles, for example, SINCR (DC+) and SINCR (LP) were prepared as described in Polo et al., *Proc. Natl. Acad. Sci. USA* (1999) <u>96</u>:4598-4603. Mice were immunized with 1 x 10³ to 1 x 10⁷ IU of SINCR (DC+) or SINCR (LP) replicon particles encoding for NS345a, by intramuscular injection into the tibialis anterior, followed by a booster injection of 10⁷ pfu vaccinia virus encoding NS5a at 6 weeks.

IFN-γ expression was measured by intracellular staining as described in Example 3. Administration of an increase in the number of SINCR (DC+) replicon particles encoding NS345a resulted in an increase in % of CD8+ T cells expressing IFN-γ.

EXAMPLE 8

Alphavirus replicon priming, followed by various boosting regimes.

Alphavirus replicon particles, for example, SINCR (DC+) were prepared as

described in Polo et al., *Proc. Natl. Acad. Sci. USA* (1999) <u>96</u>:4598-4603. Mice were
primed with SINCR (DC+), 1.5 x 10⁶ IU replicon particles encoding NS345a, by
intramuscular injection into the tibialis anterior, followed by a booster of either 10-100 μg
of plasmid DNA encoding for NS5a, 10¹⁰ adenovirus particles encoding NS345a, 1.5 x 10⁶
IU SINCR (DC+) replicon particles encoding NS345a, or 10⁷ pfu vaccinia virus encoding
NS5a at 6 weeks.

IFN-γ expression was measured by intracellular staining as described in Example 3. The results are shown in Table 7. The results demonstrate that boosting with vaccinia virus encoding NS5a DNA results in the strongest generation of CD8+ HCV specific T cells which express IFN-γ. Boosting with plasmid encoding NS5a DNA also results in a good response, while lesser responses are noted with adenovirus NS345a or SINCR DC+ boosted animals.

Table 7: Alphav	virus Replicon Par	ticle Prin	ing, Follow	ed by Variou	ıs Boostin	g Regimens
		(%		N-gamma that are IFN-	·g+)	
Vaccines	Boost	Mean	SdtdevP	# of mice tested	# of expts	% respond- ing mice
SINCR (DC+) 1.5X10 ⁶	NS5a DNA	0.46	0.36	4	1	75%
SINCR (DC+) 1.5X10 ⁶	Adeno NS345a (10X10 ¹⁰)	0.04	0.04	4	1	25%
SINCR (DC+) 1.5X10 ⁶	SINCR (DC+) 1.5X10 ⁶	0.06	0.06	. 8	2	25%
SINCR (DC+) 1.5X10 ⁶	VVNS5a (1X10 ⁷)	2.43	2.45	4	1	100%

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EXAMPLE 9

Alphaviruses expressing NS345a

Alphavirus replicon particles, for example, SINCR (DC+) and SINCR (LP) were prepared as described in Polo et al., *Proc. Natl. Acad. Sci. USA* (1999) 96:4598-4603. Mice were immunized with 1 x 10² to 1 x 10⁶ IU SINCR (DC+) replicons encoding NS345a via a combination of delivery routes (2/3 IM and 1/3 S/C) as well as by S/C alone, or with 1 x 10² to 1 x 10⁶ IU SINCR (LP) replicon particles encoding NS345a via a combination of delivery routes (2/3 IM and 1/3 S/C) as well as by S/C alone.

The immunizations were followed by a booster injection of 10⁷ pfu vaccinia virus encoding NS5a at 6 weeks.

IFN-γ expression was measured by intracellular staining as described in Example 3. The results are shown in Figure 6. The results indicate activation of CD8+ HCV specific T cells.

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EXAMPLE 10

Efficiency of NS5a encoding DNA vaccine formulations to prime CTLs in mice

Mice were immunized with either 10-100 μ g of plasmid DNA encoding NS5a as described in Example 1 or with PLG-linked DNA encoding NS5a as described in Example 5. The immunizations were followed by a booster injection at 6 weeks of either 10-100 μ g of plasmid DNA encoding for NS5a, 10^{10} adenovirus particles encoding NS345a, 1.5×10^6 IU SINCR (DC+) replicon particles encoding NS345a, or 10^7 pfu vaccinia virus encoding NS5a.

CTL activity and IFN- γ expression were measured by the methods described in Examples 2 and 3.

The results are shown in Table 8. The results demonstrate that priming with plasmid DNA encoding for NS5a or PLG-linked DNA encoding NS5a results in activation of CD8+ HCV specific T cells.

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\neg		5	· ·	· ·			
<u>.</u> 9		CTL activity	YES	YES	YES	ON	YES
ime CTLs in M		fold increase vs. 'naked' DNA	N/A	N/A:	1.29	N/A	N/A
ations to Pr		# of expts	3	É	2	. 1	1
cine Formul	ICS for IFN-gamma (%CD8+ cells that are IFN- g+)	% respond- ing	100%	%83%	100%	%05	%SL
ONA Vac	ICS for IFN-gamma CD8+ cells that are II g+)	# of mice tested	8	12	6	4	4
Table 8: Efficiency of NS5a-Encoding DNA Vaccine Formulations to Prime CTLs in Mice	ICS (%CD8-	SdtdevP	69.1	60.0	60:0	80.0	21.0
		Mean	1.67	0.17	0.22	0.10	0.20
e 8: Efficienc	·	Boost	VVNS5a	NS5a DNA	NS5a DNA	AdenoNS 345a	SINCRNS 345a
Tabl		NS5a Vaccines	NS5a DNA	NS5a DNA	PLGNS5a DNA	NS5a DNA	NS5a DNA

EXAMPLE 11

Efficiency of NS345b-encoding DNA vaccine formulations to prime CTLs in mice

Mice were immunized with 10-100 μg of plasmid DNA encoding NS34b by intramuscular injection to the tibialis anterior or with PLG linked DNA encoding NS5a as described in Example 5. The immunizations were followed by a booster injection of plasmid DNA encoding for NS5a as described in Example 1.

CTL activity and IFN-γ expression were measured by the methods described in Examples 2 and 3.

The results are shown in Table 9. The results demonstrate that priming with plasmid DNA encoding NS345b or PLG-linked NS345b results in activation of CD8+ HCV specific T cells.

	Table 9: Efficiency of NS345b-Encoding DNA Vaccine Formulations to Prime CTLs in Mice	of NS3451	b-Encoding DN	A Vaccine For	mulations to Pa	rime CTLs	in Mice	
,			ICS (%CD8+	ICS for IFN-gamma (%CD8+ cells thät are IFN-g+)	na FN-g+)		·	:
							fold increase	
NS345 DNA				# of mice	%	# of	vs. 'naked'	CIL
Vaccines	Boost	Mean	SdtdevP	tested	responding	expts	DNA	activity?
NS345 DNA	NS5a DNA	0.18	0.16	15	60%	3.	N/A	YES
PLGNS345 DNA	NS5a DNA	0:30	0.33	14	57%	3	1.67	YES

EXAMPLE 12

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Administration of DNA via separate plasmids

Mice were immunized with 100 μg plasmid DNA encoding for NS345a or with 100 μg PLG-linked DNA encoding NS345a. Additionally, separate DNA plasmids encoding NS5a, NS34a, and NS4ab (33.3 μg each) were administered concurrently to another group of mice. Finally, PLG-linked DNA encoding NS5a, NS34a, and NS4ab (33.3 μg each) were administered concurrently to another group of mice. The immunizations were followed by a booster injection of 1x10⁷ pfu vaccinia virus encoding NS5a, 6 weeks post first immunization.

IFN-γ expression was measured by the method described in Example 3. The results are shown in Figure 7. The results demonstrate a particularly vigorous response in the activation of CD8+ HCV specific T cells when the DNA is broken down into smaller sub units and linked to PLG.

15 **EXAMPLE 13**

Immunogenicity of NS345Core₁₂₁-ISCOMS in Mice

Groups of 10 C57 black mice were immunized IM at 0, 21 and 60 days with the formulations shown in Table 10. The NS345Core₁₂₁-PLGdss group received a vaccine dose of 50 μ l in each leg whereas the other vaccine groups received a vaccine dose of 50 μ l in one leg.

NS345Core₁₂₁-ISCOMS were comprised of amino acids 1242 to 3011 and 1-121 and the HCV polyprotein, numbered relative to HCV-1and were adsorbed to ISCOMS with a ratio of protein to QH of approximately 8:1, using standard techniques. See, e.g., International Publication No. WO 01/37869A.

Core-ISCOMS including an HCV core protein fragment from the region spanning amino acid positions 1-191 of the HCV polyprotein, numbered relative to HCV-1, with a ratio of protein to QH of 1:1, were produced using standard techniques. See, e.g., International Publication No. WO 01/37869A.

NS345Core₂₂ was formulated in 0.1% SDS in PBS and contained DTT.

Protein was diluted in PBS and mixed 1:1 with MF59 (see, Ott et al., "MF59 --

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Design and Evaluation of a Safe and Potent Adjuvant for Human Vaccines" in Vaccine Design: The Subunit and Adjuvant Approach (Powell, M.F. and Newman, M.J. eds.) Plenum Press, New York (1995) pp. 277-296; and U.S. Patent No. 6,299,884) prior to immunization.

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For NS345Core₁₂₁-PLGdss, PLG microparticles produced as described above were treated with 3-(trimethylsilyl)-1-propanesulfonic acid (DSS) to enhance adsorption of antigen. DSS is commercially available from, e.g., Sigma Chemical Co., St. Louis, MO. NS345Core₁₂₁ was adsorbed thereto using standard techniques (see, International Publication No. WO 00/050006). The NS345Core₁₂₁-PLGdss was mixed with MF59 prior to immunization.

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As shown in Table 10, NS345Core₁₂₁-ISCOMS produced antibody response only to NS5 in immunized C57 black mice. Higher levels of antibodies to NS5 were produced in mice immunized with NS345Core₁₂₁ adjuvanted with MF59, however no antibody response to core, NS3 or NS4 was produced with this adjuvant either.

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Mice immunized with Core-ISCOMS produced antibodies to core. In contrast, NS345Core₁₂₁-PLGdss immunized mice produced significantly higher antibodies to NS5 than NS345Core₁₂₁-ISCOMS. In addition, NS345Core₁₂₁-PLGdss immunized mice produced antibodies to NS3 and some antibody response to core, but no antibodies to NS4.

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Table 10. Immunogenicity of NS345Core₁₂₁-Iscoms in Mice. Geometric mean EIA antibody titers to core and nonstructural proteins are shown. The number of responding mice per group are also listed.

Vaccine	IM Protein	Anti-Core	Anti-C33C	Anti-C100	Anti-NS5
	Dose (μg) ^a	Antibody	(NS3)	(NS4)	Antibody
		EIA GMT	Antibody	Antibody	EIA GMT
			EIA GMT	EIA GMT	
NS345Core	6.0, 6.0, 6.0 ^b	<10	<10	<10	31
121 ISCOMS		(0/10)	(0/10)	(0/10)	(7/10)
Core-	6.0, 6.0, 6.0°	188	<10	<10	<10
ISCOMS		(9/10)	(1/10)	(0/10)	(2/10)
NS345Core	6.0, 6.0, 6.0	<10	<10	<10	279
₁₂₁ MF59		(2/9)	(1/9)	(0/9)	(9/)
NS345Core	10, 10, 10	5	50	<10	419
121 PLG-		(6/10)	(9/10)	(2/10)	(9/9)
dss/MF59		1			

 $^{\circ}$ Groups of 10 C57 black mice were immunized IM at 0, 21 and 60 days. Serum was obtained after the last immunization. The NS345 Core₁₂₁-PLGdss group received vaccine dose of 50 μ l in each leg whereas the other vaccine groups received vaccine dose of 50 μ l in one leg.

EXAMPLE 14

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25 Immunogenicity of Different Formulations of NS345 Core₁₂₁ or NS345 in Mice

Groups of 10 C57 black mice were immunized IM at 0, 30 and 60 days with the formulations shown in Tables 11 and 12. For the studies shown in Table 11, the NS345 and NS345Core₁₂₁ protein concentration was 10 μ g per dose, and for those in Table 12, the concentration was 5 μ g per dose.

^bThe ratio of protein to QH was approximately 8:1.

The ratio of protein to QH was approximately 1:1.

For PLG-NS345 (amino acids 1242 to 3011 of the HCV polyprotein) and PLG-NS345Core₁₂₁ (amino acids 1242-3011 and 1-121 of the HCV polyprotein), PLG microparticles were prepared and NS345 or NS345Core₁₂₁ were adsorbed thereto using standard techniques, as described above.

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For PLG-NS345 + PLG-CTAB-E1E2 DNA, PLG microparticles were prepared and NS345 was adsorbed to the microparticles as described above. E1E2 DNA was produced as follows. Mammalian expression plasmid pMH-E1E2-809 (Figure 4, ATCC Deposit No. PTA-3643) encodes an E1E2 fusion protein which includes amino acids 192-809 of HCV 1a (see, Choo et al., Proc. Natl. Acad. Sci. USA (1991) 88:2451-2455). Chinese Hamster Ovary (CHO) cells were used for expression of the HCV E1E2 sequence from pMH-E1E2-809. In particular, CHO DG44 cells were used. These cells, described by Uraub et al., Proc. Natl. Acad. Sci. USA (1980) 77:4216-4220, were derived from CHO K-1 cells and were made dihydrofolate reductase (dhfr) deficient by virtue of a double deletion in the dhfr gene. DG44 cells were transfected with pMH-E1E2-809. The transfected cells were grown in selective medium such that only those cells expressing the dhfr gene could grow (Sambrook et al., supra). Isolated CHO colonies were picked (~800 colonies) into individual wells of a 96-well plate. From the original 96-well plates, replicates were made to perform expression experiments. The replicate plates were grown until the cells made a confluent monolayer. The cells were fixed to the wells of the plate and permeablized using cold methanol. Anti-E1 and anti-E2 antibodies, 3D5C3 and 3E5-1 respectively, were used to probe the fixed cells. After adding an anti-mouse HRP conjugate, followed by substrate, the cell lines with the highest expression were determined. The highest expressing cell lines were then expanded to 24-well cluster plates. The assay for expression was repeated, and again, the highest expressing cell lines were expanded to wells of greater volume. This was repeated until the highest expressing cell lines were expanded from 6-well plates into tissue culture flasks. At this point there was sufficient quantity of cells to allow accurate count and harvest of the cells, and quantitative expression assays were done. An ELISA was performed on the cell extract, to determine high expressors.

To produce the PLG-CTAB-E1E2 DNA, PLG microparticles were treated with CTAB as described above (see, International Publication No. WO 00/050006).

For PLG-NS345Core₁₂₁ + E1E2 DNA PLG-NS345Core₁₂₁ and E1E2 DNA were produced as described above.

For PLG-NS345 or PLG-NS345Core₁₂₁ + MF59, PLG-NS345 or PLG-NS345Core₁₂₁ was combined with MF59 as described above.

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For PLG-NS345 or PLG-NS345Core₁₂₁ + CTAB-CpG, NS345 or NS345Core₁₂₁ was adsorbed to PLG as described above. The CpG molecule used was 5'-TCCATGACGTTCCTGACGTT-3' and this was treated with CTAB, as described above.

For PLG-NS345 or PLG-NS345Core₁₂₁ + QS21, the saponin adjuvant QS21 was combined with the PLG-HCV proteins.

For PLG-NS345 or PLG-NS345Core₁₂₁ + CTAB-CpG + MF59, the various components, as described above, were combined.

The remaining adjuvants used in the studies and shown in the tables are selfexplanatory.

The results of these studies are shown in Tables 11 and 12. As can be seen in Table 11, none of the formulations produced antibody responses to core, NS3 or NS4 antigens. However, PLG-NS345+CTAB-CPG in MF59 produced the highest antibody titers to NS5. PLG-NS345Core₁₂₁+QS21, PLG-NS345+CTAB-CPG, PLG-NS345Core₁₂₁+CTAB-CPG, and PLG-NS345 + QS21 produced moderate antibody titers to NS5. The other formulations produced very low antibody titers to NS5.

As can be seen in Table 12, NS345Core₁₂₁ /MF59/MPL and NS345Core₁₂₁/MF59/CpG formulations produced very high antibody titers to NS345Core₁₂₁. NS345Core₁₂₁/MF59, NS345/MF59/CpG, and NS345Core₁₂₁ /MF59/Chol/QS21 formulations produced moderate antibody titers to NS345Core₁₂₁. The other formulations produced very low or no antibody titers to NS345Core₁₂₁.

	Table 11. Immu	nogenicity of diffe	rent formulations	of HCV NS345Cor	e ₁₂₁ or NS345 in
	Mice. Geometri	c mean EIA antibo	dy titers to core an	d nonstructural pro	oteins are shown.
	Vaccine ^a	Anti-Core	Anti-C33C	Anti-C100	Anti-NS5
		Antibody EIA	(NS3)	(NS4)	Antibody EIA
		GMT	Antibody EIA	Antibody EIA	GMT
			: GMT	GMT	
5	PLG-NS345	<10	<10	<10	10
	PLG-	<10	<10	<10	15
	NS345Core ₁₂₁				
	PLG-	<10	11	<10	23
	NS345+PLG-				
10	CTAB-E1E2	·			
	DNA		į		
	PLG-	<10	<10	<10	20
	NS345Core ₁₂₁				
	+E1E2 DNA				
15	PLG-NS345	<10	<10	<10	70
	+MF59				
	PLG-	<10	<10	<10	26
	NS345Core ₁₂₁				
	+ MF59				
20	PLG-NS345 +	<10	<10	<10	350
	CTAB-CPG				
	PLG-	<10	<10	<10	271
	NS345Core ₁₂₁				
	+ CTAB-CPG				
25	PLG-NS345	<10	<10	<10	201
	+QS21		`,		
	PLG-	<10	<10	<10	505
	NS345Core ₁₂₁				
	+ QS21				

PLG-NS345 + CTAB-CPG + MF59	<10	<10	<10	1471
PLG- NS345Core ₁₂₁ + CTAB+MF59	<10	<10	<10	63

^aGroups of 10 C57 black mice were immunized IM at 0, 30 and 60 days. Serum was obtained after the last immunization. The NS345 or NS345Core₁₂₁ protein concentration was 10 μg per dose.

Vaccine*	Anti-NS345Core ₁₂₁ Antibody EIA GM
NS345Core ₁₂₁ /MF59	328
NS345Core ₁₂₁ /MF59/CpG	7,926
NS34a+NS5B+Core/MF59	12
NS34a+NS5B+Core/MF59/CpG	5
PLG-NS345Core ₁₂₁ /MF59	<10
PLG-NS345Core ₁₂₁ /MF59/CpG	<10
PLG-NS345Core ₁₂₁ /PLG-CpG	9
NS345Core ₁₂₁ /alum phosphate	34
NS345Core ₁₂₁ /alum phosphate//CpG	950
NS345/MF59/CpG	511
PLG-NS345/PLG/CpG	117
NS345Core 121/MF59/MPL	10,292
NS345Core ₁₂₁ /MF59/Chol/QS21	698
NS345Core 121/Alum phosphate/MPL	23

^aGroups of 10 C57 black mice were immunized IM at 0, 30 and 60 days. Serum was obtained after the last immunization. The NS345 or NS345Core₁₂₁protein concentration was 5 μg per dose.

EXAMPLE 15

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Lymphoproliferative Response of Different Formulations of NS345Core $_{121}$ or NS34A + NS35B + Core in Mice

Groups of 8 C57 black mice were immunized IM at 0, 30 and 60 days with the formulations shown in Table 13 and are as described above. Spleens were obtained after the last immunization. The NS345Core₁₂₁ protein concentration was 25 µg per dose. The NS34a, NS5b and core doses were 3 µg each.

The results of this study are shown in Table 13. As can be seen,

NS345Core₁₂₁/Alum/CpG, PLG-NS345Core₁₂₁/PLG/CpG, NS34a+NS5B+

Core/MF59/CpG and PLG-NS345Core₁₂₁/MF59/CpG formulations demonstrated strong LPA responses to NS5, NS34 and core antigens. The NS345Core₁₂₁/MF59 formulation also produced a strong LPA response to NS5 and NS34. Core was not tested.

Moderate LPA responses were observed to NS5, NS34 and Core antigens with PLGNS345Core₁₂₁/MF59 and NS34a +NS5B + Core/MF59 formulations. The
NS345Core₁₂₁/MF59/CpG formulation may not have been administered properly in
that no LPA response was observed in this experiment. In a subsequent experiment
as shown in Table 14, an LPA was observed to this formulation.

Groups of 8 C57 black mice were immunized once IM with the formulations shown in Table 14, produced as described above. Draining lymph nodes were obtained.

The NS345Core₁₂₁ protein concentration was 25 µg per dose.

The results of this study are shown in Table 14. As can be seen in Table 14, all the formulations tested produced a strong LPA response to NS5, NS34 and Core as well as the NS345Core₁₂₁.

Table 13. Lymphoproliferative response of different formulations of HCV NS345Core₁₂₁ or NS34A+NS5B+Core in Mice. LPA responses (cpm) to core and nonstructural proteins are shown. The number of mice in each group responding is also indicated in parentheses.

	puron	iuicoco.		******
				HIV-2 env
		SOD-C200	SOD-C22-3	(background
Vaccine ^a	SOD-NS5	(NS34)	(Core)	control)
NS345Core ₁₂₁ /MF59	2250	1800	ND	144
	(6/8)	(4/8)		
NS345Core ₁₂₁ /MF59/CpG	80	80	ND	138
PLG-NS345Core ₁₂₁ /MF59	560	120	510	93
	(2/8)	(2/8)	(2/8)	
PLG-	1600	1500	620	75
NS345Core ₁₂₁ /MF59/CpG	(6/8)	(6/8)	(8/8)	
NS34a+NS5B+Core/MF59	564	710	265	76
	(8/8)	(8/8)	(8/8)	
NS34a+NS5B+	1523	885	446	67
Core/MF59/CpG	(8/8)	(8/8)	(6/8)	
PLG-	3675	2860	370	88
NS345Core ₁₂₁ /PLG/CpG	(8/8)	(8/8)	(8/8)	
NS345Core ₁₂₁ /Alum/CpG	8450	7940	1040	82
	(8/8)	(8/8)	(6/8)	

 $^{\circ}$ Groups of 8 C57 black mice were immunized IM at 0, 30 and 60 days. Spleens were obtained after the last immunization. The NS345Core₁₂₁ protein concentration was 25 µg per dose. The NS34a, NS5B and Core doses were 3 µg each.

Table 14. Lymphoproliferative response of different formulations of HCV NS345Core ₁₂₁ in Mice. The LPA responses (cpm) form an average of three consecutive experiments to core and nonstructural proteins are shown.	ponse of different for three consecutive exp	ponse of different formulations of HCV NS345Core ₁₂₁ in Mice. The LPA refure consecutive experiments to core and nonstructural proteins are shown.	45Core ₁₂₁ in Mice. Ti instructural proteins a	he LPA responses (cr re shown.	om) form an average of
Vaccine	SOD-NS5	SOD-C200 (NS34)	SOD-C22-3 (Core)	NS345Core ₁₂₁	HIV-2 env (background control)
NS345Core ₁₂₁ /MF59	8900	8300	3000	20900	068
NS345Core ₁₂₁ /MF59/CpG	1890	1628	1200	20100	623
PLG-NS345Core _{L21} /MF59	10700	12900	1800	23700	818
PLG-NS345Core ₁₂₁ /MF59/CpG	3,600	4690	0991	27300	911
PLG-NS345Core ₁₂₁	4500	4660	09L	18150	315
PLG-NS345Core ₁₂₁ /PLG/CpG	7750	5300	1250	22980	450
NS345Core ₁₂₁ /Alum	3050	3725	744	11300	390
NS345Core ₁₂₁ /Alum/CpG	4130	4670	009	20660	480

⁴Groups of 8 C57 black mice were immunized once IM. Draining lymph nodes were obtained. The NS345Core₁₂₁ protein concentration was 25 μg per dose.

EXAMPLE 16

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Immunogenicity of Recombinant HCV Protein Vaccines Adjuvanted with ISCOMS in Rhesus Macaques

The safety and immunogenicity of HCV proteins completed with the adjuvant, Iscomatrix, was studied in Rhesus macaques. Three groups made up of four animals each were immunized IM as detailed below at week 0, 4 and 8 weeks. Vaccines were prepared as described above. The ISCOMS used lacked QH-A.

10	Group Number	n	Vaccine	Delivery
	1	4	Core-ISCOM (50 µg in 1 ml)	0.5 ml R Leg 0.5 ml L Leg
	2	4	NS345Core ₁₂₁ -ISCOM (1 mg in 1 ml)	0.5 ml R Leg 0.5 ml L Leg
	3	4	Core-ISCOM (25 µg in 0.5 ml) and NS5b-ISCOM (50 µg in 0.35 ml)	0.5 ml Core-ISCOM R Leg 0.35 ml NS5b-ISCOM L Leg

Bleeds occurred as follows and immunogenicity was determined by CTL assays, lymphoproliferation assays, FACS analysis and antibody response a previously described (Palakos, et al. (2001) *J. of Immunology* 166:3589).

Week	Bleed date	Immunized
-10		
-1	Х	
0		Х
2	Х	
4		⁵ X
6	X	
8		X
10	X	

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The immunogenicity of the different HCV recombinant protein vaccines is shown in Tables 15-17.

Table 15. The Immunogenicity of HCV Core-ISCOMS vaccine two weeks post 2nd immunization and post 3nd immunization as assessed by CTL assays, CD8+ FACS analysis, LPA stimulation index and CD4+ FACS analysis

		. 2 weeks post 3°											
		CI	D8+ ICS	(CTL)		CD4+ ICS (LPA SI)							
Macaqu e#	С	NS 3	NS4	NS5a	NS5b	С	NS3	NS4	NS5a	NS5b			
X020	- (-)					+(-)							
N001	- (-)					+(-)							
N086	- (-)					+(-)							
X010	- (-)					+(11)							

		2 weeks post 2°											
Macaqu		CI	D8+ ICS	(CTL)		CD4+ ICS (LPA SI)							
e#	С	NS 3	NS4	NS5a	NS5b	С	NS3	NS4	NS5a	NS5b			
X020	- (-)					+/-(-)							
N001	- (-)				· · ·	-(8)							
N086	- (-)					+(-)							
X010	- (-)					+/- (12)							

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Table 16. The Immunogenicity of HCV NS345Core₁₂₁-ISCOMS vaccine two weeks post 2nd immunization and post 3rd immunization as assessed by CTL assays, CD8+ FACS analysis, LPA stimulation index and CD4+ FACS analysis

2 weeks post 3° CD8+ICS (CTL) CD4+ ICS (LPA SI) Macaqu C NS3 NS4 NS5a NS5b C NS3 NS4 NS5a NS5b e# X016 - (-) +(+) -(-) -(-) -(-) -(-) +(-) -(-) +/-(-) +/-(-) X008 - (-) -(-) -(-) -(-) -(-) -(-) -(-) -(-) -(5) -(-) X021 - (-) -(-) -(-) -(-) -(-) -(-) -(-) -(-) -(-) -(-) +/- (-) -(-) +/-(-X023 -(-) -(-) +/-(-) -(-) -(-) -(-) -(-)

15 2 weeks post 2° Macagu CD8+ICS (CTL) CD4+ ICS (LPA SI) e# C NS3 NS4 NS5a NS5b С NS3 NS4 NS5a NS5b X016 - (-) +(+) -(-) +(+) +(+) -(-) +(-) +/-(-) +(-) +(-) +(-) X008 +(+) -(-) +(+) +(+) -(-) +(-) -(-) +-(-) +(-) 20 X021 - (-) +/-(-) +/-(-) -(-) -(-) +/-(-) -(-) -(-) -(-) -(-) +/-X023 +(+) +/-(-) +(-) -(-) +(-) -(-) -(-) -(-) 7 (+)

Table 17. The Immunogenicity of HCV Core-ISCOMS + NS5b-ISCOMS vaccine two weeks post 2nd immunization and post 3rd immunization as assessed by CTL assays, CD8+ FACS analysis, LPA stimulation index and CD4+ FACS analysis

	2 weeks post 3°											
		CI	D8+ ICS	(CTL)		CD4+ ICS (LPA SI)						
Macaqu e#	С	NS 3	NS4	NS5a	NS5b	С	NS3	NS4	NS5a	NS5b		
X022	+(-)				-(-)	-(8)				+/-(11)		
X014	-(-)				-(-)	+(6)				+(11)		
N154	-(-)				-(-)	-(-)				+(-)		
N173	-(-)				-(-)	-(-)				+/-(-)		
	1 .				1	1		1	Ì			

	2 weeks post 2°											
Macaqu		CI	08+ ICS	(CTL)		CD4+ ICS (LPA SI)						
e#	С	NS 3	NS4	NS5a	NS5b	С	NS3	NS4	NS5a	NS5b		
X022	-(-)				-(+)	-(-)				-(-)		
X014	+(-)				+/-(-)	-(-)				+(-)		
N154	-(-)				-(-)	-(6)				+(8)		
N173	-(-)				-(-)	+/-(-)				+(6)		

As can be seen in Table 15, the HCV Core-ISCOM vaccine produced no CTL positive responses in any of the 4 immunized macaques after the second or third immunizations. No positive CD8 γ-interferon and/or TNF-α intracellular staining was also observed, although backgrounds were high in these particular arrays. At least two of four macaques produced a strong LPA response after the second immunizations, but only one remained positive after the third immunization. Two of four macaques produced positive CD4 intracellular staining after the second immunization and four of four after the third immunization.

As shown in Table 16, the HCV NS345Core₁₂₁-ISCOM vaccine after the second immunization produced CTL positive responses to peptide pools representing two or more HCV proteins in three of four macaques (two of these macaques had responses to peptide pools from NS3, NS5a and NS5b, one to peptide pools from core and NS3). CD8 positive γ-interferon and/or TNF-α intracellular staining to peptide pools representing two or more HCV proteins was positive in at least three of four macaques. One of four macaques produced a strong LPA response. At least three of four macaques produced CD4 positive intracellular staining to two or more HCV proteins. After the third immunization, only one of four macaques had a positive CTL response, CD8 positive intracellular staining and C04 positive intracellular staining. One other macaque had a positive LPA response and weak CD8+ CD4 intracellular staining, This decline in immunogenicity was likely due to instability of the vaccine formulation (see below).

As shown in Table 17, the HCV Core-ISCOM + NS5b-ISCOM vaccine produced a CTL positive response to NS5b in one of the 4 immunized macaques after the second immunization which did not remain positive after the third immunization. CD8 positive intracellular positive staining was observed in one of four animals post second. Two of four macaques produced a strong LPA response after the second immunization which did not remain positive after the third immunization. Two other macaques did develop a strong LPA response after the third immunization. Three or four developed positive CD4 intracellular staining. One developed positive CD8 intracellular staining.

Three weeks after the third immunization, it was noted that the physical appearance of the polyprotein vaccine solution was visibly turbid. The core vaccine also was turbid but less so. The Core-NS5 vaccine was also slightly turbid. Analysis of this turbidity in the polyprotein formulation indicated that the ISCOM particles had precipitated into large aggregates. These aggregates could be dispersed by vortexing with 0.1% TWEEN 80 detergent. It is probable that this change in the formulation of the vaccine occurred before the last immunization. This observed change in appearance of the vaccines may have affected their immunogencity as cellular immune results declined in all three vaccines.

The immunogenicity of HCV Core-ISCOMS, NS345Core₁₂₁-ISCOMS and Core-ISCOMS + NS5b-ISCOMS as assessed by EIA antibody response is shown in Table 18. As can be seen, all three vaccines produced an antibody response by the third immunization to their corresponding HCV proteins, except for the NS345Core₁₂₁-ISCOM vaccine. The NS345Core₁₂₁-ISCOM vaccine produced antibody responses to NS3, NS4 and a very strong antibody response to NS5, but no antibody response to HCV core.

Table 18. The immunogenicity of HCV Core-ISCOMS, NS345Core_{tal}-ISCOMS, Core-ISOCMS + NS5b-ISCOMS vaccine two weeks post 2nd immunization and post 3rd immunization as assessed by EIA antibody response to HCV proteins.

,	IIIIIIUUIZ	auon as as	sessed by	EIA antibo	rdy respon	ise to nev	proteins.	
Vaccine Macaque		ore EIA ody Titer		S3 EIA dy Titer	Anti-N Antibo	S4 EIA dy Titer		S5 EIA dy Titer
#	Post 2 nd	Post 3 rd	Post 2 nd	Post 3 rd	Post 2 nd	Post3 rd	Post 2 nd	Post 3 rd
Core- ISCOM								
X020	66	226						
N001	87	46						
N086	363	396		·				
X010	108	137						
NS345 Core121/ ISCOM								
X016	<10	<10	<10	554:	56	68	3,590	3,405
X008	<10	<10	66	995	14	44	2,109	3,213
X021	<10	<10	128	6,330	41	204	7,213	8,083
X023	<10	<10	<10	3,910	64	64	1,243	4,704
Core- ISCOM + NS5b- ISCOM								
X022	<10	18					<10	134
X014	<10	13		:			<10	693
N154	542	554		:			<10	272
N173	28	78					<10	258

EXAMPLE 17

Immunization of Chimpanzees with Recombinant HCV Protein and DNA Vaccines

Five groups of five chimps each were immunized IM at 0, 0.7, 2 and 5

5 months

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with the formulations presented below. Blood was collected at week 0, two weeks subsequent to the second immunization, two weeks following the third immunization and two weeks after the fourth immunization.

Formulation 1: 20 μg E1E2 polypeptide + MF59 + 500 μg CpG (produced as described above);

Formulation 2: 1 mg NS345Core₁₂₁-ISCOM (produced as described above);

Formulation 3: 6 mg each of CTAB-PLG-E1E2 (bp 574-2427, encoding amino acids 192-809 of the HCV polyprotein, numbered relative to HCV-1); CTAB-PLG-NS34a (bp 3079-5133, encoding amino acids 1027-1711 of the HCV polyprotein, numbered relative to HCV-1); CTAB-PLG-NS34ab (bp 4972-5916, encoding amino acids 1658-1972 of the HCV polyprotein, numbered relative to HCV-1); CTAB-PLG-NS5a (bp 5917-7260, encoding amino acids 1973-2420 of the HCV polyprotein, numbered relative to HCV-1);

Formulation 4: 6 mg each of E1E2 DNA, NS34a DNA, NS34ab DNA and NS5a DNA, having the same coordinates as described above, delivered without PLG via electroporation (see, e.g., U.S. Patent Nos. 6,132,419; 6,451,002, 6,418,341, 6233,483, U.S. Patent Publication No. 2002/0146831; and International Publication No. WO/0045823, for this delivery technique). Results are shown in Figures 8-10.

As can be seen, in Figure 8, all vaccines were capable of priming CD4+ and CD8+ cells specific to HCV. Thus, all vaccines were successful at inducing a T cell response to HCV. Determination of the results for the PLG-DNA from formulation 3 at two weeks subsequent to the fourth vaccination is in progress.

As shown in Figures 9 and 10, multiple T cell specificities were induced by the two vaccines. Both vaccines primed T-cells specific for multiple T cell epitopes.

As can be seen in Tables 19 and 20, E1E2 adjuvanted with MF59 primed

anti-E1E2 titers. CpG enhanced anti-E1E2 responses as well as TH1 responses and the ISCOM and the two DNA vaccines were capable of priming CD4+ and CD8+ T cell responses to HCV.

E1E2NS345a or	Post 4th	6	10	50	207	358	10	•	-	•	1
roporated DNA I	Post 3 rd	•	•	198	1,197	1,267	30	309	12	29	12
nized with Elect	Post 2 nd	•	_	457	1,261	1,426	•	104	•	•	•
in chimps immu	Pre 1st	•	•	10	206	245	•	•		2	1
IA antibody titers	Chimp	4X0330	4X0335	4X0348°	4X0354 ^d	4X0368°	4X0238	4X0239	4X0250	4X0278	4X0288
Table 19. Anti-E1E2 EIA antibody titers in chimps immunized with Electroporated DNA E1E2NS345a or PLG DNA E1E2NS345	Vaccine	Electroporated DNA	E1E2-NS34A-	NS4AB-NS5A [®]			PLG DNA	E1E2-NS34A-	NS4AB-NS5A ^b		

Electroporated IM with 1.5 mg of each plasmid at 0, 0.7, 2 and 5 months.

Bleeds were taken 14 days after each immunization.

^bIM immunization with 1.5 mg of each PLG plasmid at 0, 0.7, 2 and 6 months.

Bleeds were taken 14 days after each immunization. Prior E2 immunization Prior E1E2 immunization

CD4+ Table 20. Immunogenicity in chimps of low dose (20 μ g) HCV E1E2 antigen using MF59 or MF59 combined with CpG as adjuvants (2 wks post 3^{rd}) 7 ‡ E1E2 EIA Ab GMT $P = 0.029^b$ 2,713 261 Ab Titer E1E2EIA 8,835 2,713 1,238 2,580 3,201 510 131 101 421 84 4x0346 4x0426 4x0365 4x0419 4x0410 4x0367 4x0420 4x0372 4x0431 4x0371 Chimp MF59/CpG Vaccinea E1E2/ MF59 E1E2/

*Chimps immunized IM at 0, 1 and 6 mos with 20 µg of E1E2 antigen using MF59 with or without 500 µg of CpG. Serum samples were obtained 14 days after last immunization.

^bChimps immunized with E1E2 using CpG combined with MF59 as adjuvant produced significantly higher (P<0.05) levels of E1E2 EIA antibody than chimpanzees with E1E2 using MF59 alone.*

Thus, HCV polypeptides and polynucleotides, either alone or as fusions, to stimulate cell-mediated immune responses, are disclosed. Although preferred embodiments of the subject invention have been described in some detail, it is understood that obvious variations can be made without departing from the spirit and the scope of the invention as defined by the appended claims.

We claim:

A fusion protein comprising HCV polypeptides, wherein the HCV polypeptides consist essentially of an NS3, an NS4, an NS5 and a core polypeptide of
 a hepatitis C virus (HCV), wherein said core polypeptide consists of amino acids 1 121 of the HCV polyprotein, numbered relative to the full-length HCV-1 polyprotein.

- The fusion protein of claim 1, wherein the NS5 polypeptide is an NS5a
- 10 polypeptide.
 - The fusion protein of claim 1, wherein the NS5 polypeptide is an NS5b

polypeptide.

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4. The fusion protein of claim 1, wherein the NS5 polypeptide is an NS5a

and an NS5b polypeptide.

20 6. The fusion protein of claim 4, wherein the protein comprises the sequence

of amino acids of SEQ ID NO:8.

- 7. A fusion protein according to any of claims 1-6, wherein at least one of the HCV polypeptides is derived from a different strain of HCV than the other HCV polypeptides.
 - 8. A composition comprising:
 - (a) a fusion protein according to any of claims 1-7; and
- 30 (b) a pharmaceutically acceptable excipient.

9.	The composition of claim	8, further	comprising an	adjuvant.
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10. The composition of claim 8, further comprising a CpG oligonucleotide.

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The composition of claim 8, wherein said fusion protein is adsorbed to 11.

entrapped within a microparticle or ISCOM.

- 10 The composition of claim 8, further comprising a polynucleotide 12. encoding an HCV E1E2 complex.
 - 13. An isolated and purified polynucleotide that encodes a fusion protein according to any of claims 1-7.

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- 14. A composition comprising:
 - (a) the isolated and purified polynucleotide of claim 13; and
 - (b) a pharmaceutically acceptable excipient.
- 20 15. The composition of claim 14, further comprising an adjuvant.
 - 16. The composition of claim 14, wherein said polynucleotide is adsorbed to or entrapped within a microparticle.

- 17. The composition of claim 14, further comprising a polynucleotide encoding an HCV E1E2 complex.
- A method of activating T cells of a vertebrate subject which recognize 18. 30 an epitope of an HCV polypeptide, comprising the step of: administering the composition of any of claims 8-12 to said vertebrate

subject, whereby a population of activated T cells recognizes an epitope of the NS3, NS4, NS5 and/or core polypeptides.

19. A method of activating T cells of a vertebrate subject which recognize
 5 an epitope of an HCV polypeptide, comprising the step of:

administering the composition of any of claims 14-17 to said vertebrate subject, whereby a population of activated T cells recognizes an epitope of the NS3, NS4, NS5 and/or core polypeptides.

- 10 20. The method of claim 19, wherein the polynucleotide is administered via electroporation.
- 21. Use of a composition according to any of claims 8-12 and 14-17 for activating T cells of a vertebrate subject which recognize an epitope of an HCV polypeptide, wherein a population of activated T cells recognizes an epitope of the NS3, NS4, NS5 and/or core polypeptides.
- 22. Use of a fusion protein according to any of claims 1-7 for the
 manufacture of a medicament for activating T cells of a vertebrate subject which
 recognize an epitope of an HCV polypeptide, wherein a population of activated T
 cells recognizes an epitope of the NS3, NS4, NS5 and/or core polypeptides.
- Use of a polynucleotide according to claim 13 for the manufacture of a

medicament for activating T cells of a vertebrate subject which recognize an epitope of an HCV polypeptide, wherein a population of activated T cells recognizes an epitope of the NS3, NS4, NS5 and/or core polypeptides.

HCV Genome and Recombinant Proteins

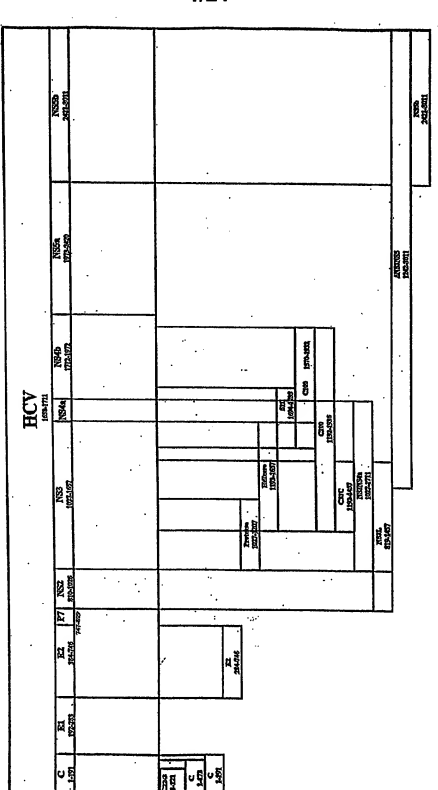


FIG 1

					M ATG	A GCG	ccc P	I ATC	T ACG	A GCG	Y TAC	A GCC	Q CAG	Q CAG
T ACA	R AGG	GGC	L CTC	L CTA	GGG G	C TGC	I ATA	·I	20 T. ACC	g Agc	L CTA	T ACT	G G	R CGG
D GAC	K AAA	n Aac	Q CAA-	GIG A	E GAG	g GGT	E GAG	V GTC	Q CAG	I ATT	V GTG	S TCA	T ACT	40 A GCT
A GCC	CAA Q	T ACC	F TTC	L CTG	A GCA	T ACG	C TGC	I ATC	50 N AAT	GGG G	V GTG	C TGC	TGG W	T ACT
V GTC	Y TAC	H CAC	G G GGG	A GCC	G GGA	T ACG	r Agg	T ACC	I ATC	A. GCG	s TCA	P CCC	K AAG	70 G GGT
P CCT	V GTC	I ATC	Q CAG	M ATG	Y TAT	T ACC	n TÄA	y GTA	BO D GAC	Q CAA	D GAC	L CTT	gig V	G GGC
w T G G	CCC P	A GCT	P CCG	90 CAA CAA	G GGT	g . Agc	R CGA	s TCA	L TTG	T ACA	CCC	C	T ACT	· c
G G G	s TCC	g TCG	D GAC	L CTT	Y TAC	L CTG	V GTC	T	110 R AGG	H CAC	A GCC	D GAT	gtc V	i AŢT
CCC E	V GTG	R CGC	R CGG	120 R CGG	ggt G	'D GAT	8 AGC	r agg	G GGC	s AGC	L CTG	L CTG	e TCG	130 P CCC
R CGG	. CCC	Į ATT	s TCC	Y TAC	L TTG	K AAA	G	s TCC	140 S TCG	GGG G	g Ggt	CCG p	L CTG	L TTG
C TGC	CCC	A GCG	GGG G	150 H CAC	A GCC	QTG V	GGC G	I ATA	. F TTT	R AGG	A GCC	A GCG	gig V	160 C TGC
T ACC	r Cgt	G GGA	V GTG	a GCT	K RAG	A GCG	V GTG	D GAC	170 F TTT	I ATC	P CCT	GIG A	E ·	n aac
L CTA	E GAG	T ACA	T ACC	180 M ATG	r Agg	s TCC								

MATURE E1	
SerPheSerIlePheLeuLeuAlaLeuLeuSerCysLeuThrValProAlaSerAlaTyr	192
TCTTTCTCTATCTTCCTTCTGGCCCTGCTCTCTTGCTTGACTGTGCCCGCTTCGGCCTAC	
AGAAAGAGATAGAAGGAAGACCGGGACGAGAGAACCGAACTGACACGGGCGAAGCCGGATG	
GlnValArgAsnSerThrGlyLeuTyrHisValThrAsnAspCysProAsnSerSerIle	212
CAAGTGCGCAACTCCACGGGGCTCTACCACGTCACCAATGATTGCCCTAACTCGAGTATT	
GTTCACGCGTTGAGGTGCCCCGAGATGGTGCAGTGGTTACTAACGGGATTGAGCTCATAA	
ValTyrGluAlaAlaAspAlaIleLeuHisThrProGlyCysValProCysValArgGlu	232
GTGTACGAGGCGGCCGATGCCATCCTGCACACTCCGGGGTGCGTCCCTTGCGTTCGCGAG	
$\tt CACATGCTCCGCCGGCTACGGTAGGACGTGTGAGGCCCCCACGCAGGGAACGCAAGCGCTC$	
GlyAsnAlaSerArgCysTrpValAlaMetThrProThrValAlaThrArgAspGlyLys	
GGCAACGCCTCGAGGTGTTGGGTGGCGATGACCCCTACGGTGGCCACCAGGGATGGCAAA	252
GOCARCGCCTCGAGGTGTTGGGTGGCGATGACCCCTACGGTGGCCACCAGGGATGGCAAA CCGTTGCGGAGCTCCACAACCCACCGCTACTGGGGATGCCACCGGTGGTCCCTACCGTTT	
CCGTTGCGGAGCTCCACACCCACCGCTACTGGGGATGCCACCGGTGGTCCCTACCGTTT	
LeuProAlaThrGlnLeuArgArgHisIleAspLeuLeuValGlySerAlaThrLeuCys	272
CTCCCCGCGACGCAGCTTCGACGTCACATCGATCTGCTTGTCGGGAGCGCCACCCTCTGT	
GAGGGGCGCTGCGTCGAAGCTGCAGTGTAGCTAGACGAACAGCCCTCGCGGTGGGAGACA	
SerAlaLeuTyrValGlyAspLeuCysGlySerValPheLeuValGlyGlnLeuPheThr	292
TCGGCCCTCTACGTGGGGGACCTGTGGGGGGTCTGTCTTTCTT	
AGCCGGGAGATGCACCCCTGGACACGCCCAGACAGAAAGAA	
PheSerProArgArgHisTrpThrThrGlnGlyCysAsnCysSerIleTyrProGlyHis	312
TTCTCTCCCAGGCGCCACTGGACGACGCAAGGTTGCAATTGCTCTATCTA	
AAGAGAGGGTCCGCGGTGACCTGCTGCGTTCCAACGTTAACGAGATAGAT	
${\tt IleThrGlyHisArgMetAlaTrpAspMetMetMetAsnTrpSerProThrThrAlaLeu}$	332
ATAACGGGTCACCGCATGGCATGGGATATGATGAACTGGTCCCCTACGACGGCGTTG	
${\tt TATTGCCCAGTGGCGTACCGTACCTATACTACTTGACCAGGGGATGCTGCCGCAAC}$	
ValMetAlaGlnLeuLeuArgIleProGlnAlaIleLeuAspMetIleAlaGlyAlaHis	252
GTAATGGCTCAGCTGCTCCGGATCCCACAAGCCATCTTGGACATGATCGCTGGTGCTCAC	354
CATTACCGAGTCGACGAGGCCTAGGGTGTTCGGTAGAACCTGTACTAGCGACCACGAGTG	
CATTACCGAGTCGACGAGGCTAGGGTGTTCGGTAGTACTAGCGACCACGAGTG	
TrpGlyValLeuAlaGlyIleAlaTyrPheSerMetValGlyAsnTrpAlaLysValLeu	372
TGGGGAGTCCTGGCGGGCATAGCGTATTTCTCCATGGTGGGGAACTGGGCGAAGGTCCTG	
${\tt ACCCCTCAGGACCGCCCGTATCGCATAAAGAGGTACCACCCCTTGACCCGCTTCCAGGAC}$	
EQ.	
E2 ValValLeuLeuLeuPheAlaGlyValAspAlaGluThrHisValThrGlyGlySerAla	302
GTAGTGCTGCTGTTTTTGCCGGCGTCGACGCGGAAACCCACGTCACCGGGGGAAGTGCC	J 7 6
CATCACGACGATAAACGGCCGCAGCTGCGCCTTTGGGTGCAGTGGCCCCCTTCACGG	
- TOTOLOGIONALIMACOGCOCHOCIOCOCII I I I I I I I I I I I I I I I	
GlyHisThrValSerGlyPheValSerLeuLeuAlaProGlyAlaLysGlnAsnValGln	412
GGCCACACTGTGTCTGGATTTGTTAGCCTCCTCGCACCAGGCGCCCAAGCAGAACGTCCAG	

	CTGATCAACACCAACGGCAGTTGGCACCTCAATAGCACGGCCCTGAACTGCAATGATAGCGCCTAGTTGTGTTGCCGTCAACCGTGGAGTTATCGTGCCGGGACTTGACGTTACTATCG	
	LeuAsnThrGlyTrpLeuAlaGlyLeuPheTyrHisHisLysPheAsnSerSerGlyCys CTCAACACCGGCTGGTTGGCAGGGCTTTTCTATCACCACAAGTTCAACTCTTCAGGCTGT GAGTTGTGGCCGACCAACCGTCCCGAAAAGATAGTGGTGTTCAAGTTGAGAAGTCCGACA	
	ProgluargleualaserCysargProLeuThrAspPheAspGlnGlyTrpGlyProIle CCTGAGAGGCTAGCCAGCTGCCGACCCCTTACCGATTTTGACCAGGGCTGGGGCCCTATC GGACTCTCCGATCGGTCGACGGCTGGGGAATGGCTAAAACTGGTCCCGACCCCGGGATAG	
	SerTyrAlaAsnGlySerGlyProAspGlnArgProTyrCysTrpHisTyrProProLys AGTTATGCCAACGGAAGCGGCCCCGACCAGCGCCCCTACTGCTGGCACTACCCCCCAAAA TCAATACGGTTGCCTTCGCCGGGGCTGGTCGCGGGGATGACGACCGTGATGGGGGGGTTTT	
	ProCysGlyIleValProAlaLysSerValCysGlyProValTyrCysPheThrProSer CCTTGCGGTATTGTGCCCGCGAAGAGTGTGTGTGTGCGGTATATTGCTTCACTCCCAGC GGAACGCCATAACACGGGCGCTTCTCACACACACAGGCCATATAACGAAGTGAGGGTCG	512
•	ProValValValGlyThrThrAspArgSerGlyAlaProThrTyrSerTrpGlyGluAsn CCCGTGGTGGTGGGAACGACCGACAGGTCGGGCGCGCCCCCCCTACAGCTGGGGTGAAAAT GGGCACCACCCTTGCTGGCTGTCCAGCCCGCGGGGTGGATGTCGACCCCACTTTTA	532
	AspThraspValPheValLeuAsnAsnThrArgProProLeuGlyAsnTrpPheGlyCys GATACGGACGTCTTCGTCCTTAACAATACCAGGCCACCGCTGGGCAATTGGTTCGGTTGT CTATGCCTGCAGAAGCAGGAATTGTTATGGTCCGGTGGCGACCCGTTAACCAAGCCAACA	552
	thm:thm:thm:thm:thm:thm:thm:thm:thm:thm:	572
	GlyAlaGlyAsnAsnThrLeuHisCysProThrAspCysPheArgLysHisProAspAla GGGGCGGGCAACACACCCTGCACTGCCCCCACTGATTGCTTCCGCAAGCATCCGGACGCC CCCCGCCCGTTGTTGTGGGACGTGACGGGGTGACTAACGAAGGCGTTCGTAGGCCTGCGG	592
	ThrTyrSerArgCysGlySerGlyProTrplleThrProArgCysLeuValAspTyrPro ACATACTCTCGGTGCGGCTCCGGTCCCTGGATCACACCCAGGTGCCTGGTCGACTACCCG TGTATGAGAGCCACGCCGAGGCCAGGGACCTAGTGTGGGTCCACGGACCAGCTGATGGGC	612
	TyrArgLeuTrpHisTyrProCysThrlleAsnTyrThrllePheLysIleArgMetTyr TATAGGCTTTGGCATTATCCTTGTACCATCAACTACACTATATTTAAAATCAGGATGTAC ATATCCGAAACCGTAATAGGAACATGGTAGTTGATGTGATATAAATTTTAGTCCTACATG	632
	ValGlyGlyValGluHisArgLeuGluAlaAlaCysAsnTrpThrArgGlyGluArgCys GTGGGAGGGGTCGAGCACAGGCTGGAAGCTGCCTGCAACTGGACGCGGGGCGAACGTTGC CACCCTCCCCAGCTCGTGCCGACCTTCGACGGACGTTGACCTGCGCCCCGCTTGCAACG	652
	AspleuGluAspArgAspArgSerGluLeuSerProLeuLeuLeuThrThrThrGlnTrp GATCTGGAAGATAGGGACAGGTCCGAGCTCAGCCCGTTACTGCTGACCACTACACAGTGG CTAGACCTTCTATCCCTGTCCAGGCTCGAGTCGGGCAATGACGACTGGTGATGTCTCACC	672

GlnValLeuProCysserPheThrThrLeuProAlaLeuSerThrGlyLeuIleHisLeu CAGGTCCTCCCGTGTTCCTTCACAACCCTGCCAGCCTTGTCCACCGGCCTCATCCACCTC GTCCAGGAGGGCACAAGGAAGTGTTGGGACGGTCGGAACAGGTGGCCGGAGTAGGTGGAG	692
HisGlnAsnIleValAspValGlnTyrLeuTyrGlyValGlySerSerIleAlaSerTrpCACCAGAACATTGTGGACGTGCAGTACTTGTACGGGGTGGGGTCAAGCATCGCGTCCTGGGTGGG	712
AlaIleLysTrpGluTyrValValLeuLeuPheLeuLeuAlaAspAlaArgValCys GCCATTAAGTGGGAGTACGTCGTCCTCCTGTTCCTTCTGCTTGCAGACGCGCGCG	732
P7	
SerCysLeuTrpMetMetLeuLeuIleSerGlnAlaGluAlaAlaLeuGluAsnLeuVal TCCTGCTTGTGGATGATGCTACTCATATCCCAAGCGGAAGCGGCTTTGGAGAACCTCGTA AGGACGAACACCTACTACGATGAGTATAGGGTTCGCCTTCGCCGAAACCTCTTGGAGCAT	
IleLeuAsnalaalaserLeuAlaGlyThrHisGlyLeuValSerPheLeuValPhePhe ATACTTAATGCAGCATCCCTGGCCGGGACGCCCGGTCTTGTATCCTTCCT	
CysPheAlaTrpTyrLeuLysGlyLysTrpValProGlyAlaValTyrThrPheTyrGly TGCTTTGCATGGTATCTGAAGGGTAAGTGGGTGCCCGGAGCGGTCTACACCTTCTACGGG ACGAAACGTACCATAGACTTCCCATTCACCCACGGCCTCGCCAGATGTGGAAGATGCCC	
MetTrpProLeuLeuLeuLeuLeuAlaLeuProGlnArgAlaTyrAlaOC ATGTGGCCTCTCCTGCTGCTGTTGGCGTTGCCCCAGCGGGCGTACGCGTAA TACACCGGAGAGGAGGACGAGGACAACCGCAACGGGGTCGCCCGCATGCGCATT	809

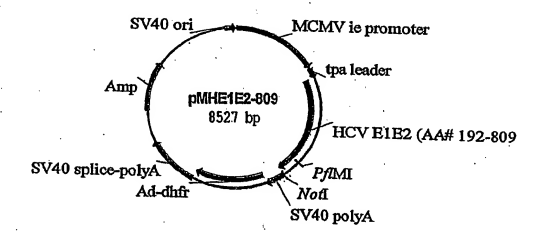


FIG. 4

7/21

M A A Y A A Q G Y K ATG GCT GCA TAT GCA GCT CAG GGC TAT AAG

V L V L N P S V A A T L G F GTG CTA GTA CTC AAC CCC TCT GTT GCT GCA ACA CTG GGC TTT GGT. 30 S K A H G I D P N I R Y M GCT TAC ATG TCC AAG, GCT CAT GGG ATC GAT CCT AAC ATC AGG ACC ..G V R T I T T G S P I T Y S GGG GTG AGA ACA ATT ACC ACT GGC AGC CCC ATC ACG TAC TCC ACC 60 F L A D G G C S G A Y TAC GGC AAG TTC CTT GCC GAC GGC GGG TGC TCG GGG GGC GCT TAT I I C D E C ·H S T GAC ATA ATA ATT TGT GAC GAG TGC CAC TCC ACG GAT GCC ACA TCC • 90 L G I G TVLDQAETA ATC TTG GGC ATT GGC ACT GTC CTT GAC CAA GCA GAG ACT GCG GGG 110 RLVVLATAT GCG AGA CTG GTT GTG CTC GCC ACC GCC ACC CCT CCG GGC TCC GTC 120 TVPHPNIEEVALSTT ACT GTG CCC CAT CCC AAC ATC GAG GAG GTT GCT CTG TCC ACC ACC 140 G E I P F Y G K A I P L GGA GAG ATC CCT TTT TAC GGC AAG GCT ATC CCC CTC GAA GTA ATC 150 160 K G G R H L I F C H S K K K C AAG GGG GGG AGA CAT CTC ATC TTC TGT CAT TCA AAG AAG AAG TGC 170 DELAAKLVALGINAV GAC GAA CTC GCC GCA AAG CTG GTC GCA TTG GGC ATC AAT GCC GTG 180 AYYRGLDVSVIPTSG

FIG. 5A

GCC TAC TAC CGC GGT CTT GAC GTG TCC GTC ATC CCG ACC AGC GGC

								A				G GGC		
												V GTC		
												E GAG		
												R CGG		
T ACT	G GGC	R AGG	G GGG	K AAG	P CCA	G GGC	I ATC	Y TAC	260 R AGA	F TTT	V GTG	A GCA	P CCG	G GGG
E GAG	R _. CGC	P	s TCC	270 G GGC	M ATG	F TTC	D GAC	S TCG	s TCC	V GTC	L CTC	C TGT	E GAG	280 C TGC
												A GCC		
T ACA	V GTT	R AGG	L CTA	300 R CGA	A GCG	Y TAC	M ATG	N. N.	T ACC	P CCG	G GGG	L CTT	P	310 V GTG
C TGC	Q CAG	D GAC	H CAT	L CTT	E GAA	F TTT	W TGG	E GAG	320 G GGC	V GTC	F TTT	T ACA	G GGC	L CTC
T ACT	H CAT	I ATA	D GAT	330 A GCC	H CAC	F TTT	L CTA	s TCC	Q CAG	T ACA	K AAG	Q CAG	S AGT	340 G GGG
E GAG	N AAC	L CTT	P CCT	Y TAC	L CTG	V GTA	A GCG	Y TAC	350 Q CAA	A GCC	T ACC	V GTG	C TGC	A GCT
												W TGG		
												P		

													H CAC	
TAC	AGA	CTG	GGC	GCT	GII	CAG	AAT	GAA	ATC	ACC	CTG	ACG	CAC	CCA
			Y TAC					M.		A GCC		L CTG	E GAG	V GTC
			T ACC										A GCT	
··· A. GCC													G GGC	
													E GAA	
				F TTC									H CAC	L TTA
													K AAG	
	A GCC			L CTC								A GCA	E GAG	
I ATC		P CCT		510 V GTC									T ACC	520 F TTC
W TGG	A GCG		H CAT			N AAC				G GGG		Q CAA	Y TAC	
A GCG	G GGC			540 T ACG									S TCA	550 L TTG
M ATG	A GCT	F TTT	T ACA	A GCT	A GCT	V GTC	T ACC	S AGC	560 P CCA	L CTA	T ACC	T ACT	S AGÇ	Q CAA
			F TTC		I ATA	L TTG	G GGG	G GGG	W TGG	V GTG	A GCT.	A GCC	Q CAG	580 L CTC

10/21

590 A A P G A A T A F V G A G L A GCC GCC CCC GGT GCC GCT ACT GCC TTT GTG GGC GCT GGC TTA GCT 600 610 G A A I G S V G L G K V L I D. GGC GCC GCC ATC GGC AGT GTT GGA CTG GGG AAG GTC CTC ATA GAC 620 I L A G Y G A G V A G A L V A ATC CTT GCA GGG TAT GGC GCG GGC GTG GCG GGA GCT CTT GTG GCA 630 F K I M S G E V P S T E D L V TTC AAG ATC ATG AGC GGT GAG GTC CCC TCC ACG GAG GAC CTG GTC 650 N L L P A I L S P G A L V V G AAT CTA CTG CCC GCC ATC CTC TCG CCC GGA GCC CTC GTA GTC GGC 660 V V C A A I L R R H V G P G E GTG GTC TGT GCA GCA ATA CTG CGC CGG CAC GTT GGC CCG GGC GAG 680 G A V Q W M N R L I A F A S R GGG GCA GTG CAG TGG ATG AAC CGG CTG ATA GCC TTC GCC TCC CGG GNHVSPTHYVPESDA 690 GGG AAC CAT GTT TCC CCC ACG CAC TAC GTG CCG GAG AGC GAT GCA 710 A A R V T A I L S S L T V T Q GCT GCC CGC GTC ACT GCC ATA CTC AGC AGC CTC ACT GTA ACC CAG 720 LLRRLHQWISSECTT CTC CTG AGG CGA CTG CAC CAG TGG ATA AGC TCG GAG TGT ACC ACT 740 PCSGSWLRDIWDWIC CCA TGC TCC GGT TCC TGG CTA AGG GAC ATC TGG GAC TGG ATA TGC 750 E V L S D F K T W L K A K L M GAG GTG TTG AGC GAC TTT AAG ACC TGG CTA AAA GCT AAG CTC ATG 770 PQLPGIPFVSCQRGY CCA CAG CTG CCT GGG ATC CCC TTT GTG TCC TGC CAG CGC GGG TAT

11/21

790 K .G V W R G D G I M H T R C H 780 AAG GGG GTC TGG CGA GGG GAC GGC ATC ATG CAC ACT CGC TGC CAC 800 C G A E I T G H V K N G T M R TGT GGA GCT GAG ATC ACT GGA CAT GTC AAA AAC GGG ACG ATG AGG 810 820 I V G P R T C R N M W S G T F 810 ATC GTC GGT CCT AGG ACC TGC AGG AAC ATG TGG AGT GGG ACC TTC $^{\circ \circ}$. $^{\circ}$ P I N A Y T T G P C T P L P A CCC ATT AAT GCC TAC ACC ACG GGC CCC TGT ACC CCC CTT CCT GCG 840 PNYTFALWRVSAEEY CCG AAC TAC ACG TTC GCG CTA TGG AGG GTG TCT GCA GAG GAA TAC 860 V E I R Q V G D F H Y V T G M GTG GAG ATA AGG CAG GTG GGG GAC TTC CAC TAC GTG ACG GGT ATG 870 880 -T T D N L K C P C Q V P S P E ACT ACT GAC AAT CTT AAA TGC CCG TGC CAG GTC CCA TCG CCC GAA 900 P C K P L L R E E V S F R V G CCC TGC AAG CCC TTG CTG CGG GAG GAG GTA TCA TTC AGA GTA GGA 920 L H E Y P V G S Q L P C E P E CTC CAC GAA TAC CCG GTA GGG TCG CAA TTA CCT TGC GAG CCC GAA 930 P D V A V L T S M L T D P S H CCG GAC GTG GCC GTG TTG ACG TCC ATG CTC ACT GAT CCC TCC CAT 950 I T A E A A G R R L A R G S P ATA ACA GCA GAG GCG GCC GGG CGA AGG TTG GCG AGG GGA TCA CCC 960 970 PSVASSSASQLSAPS CCC TCT GTG GCC AGC TCC TCG GCT AGC CAG CTA TCC GCT CCA TCT

12/21

980 L K A T C T A N H D S P D A E CTC AAG GCA ACT TGC ACC GCT AAC CAT GAC TCC CCT GAT GCT GAG 990 1000 L I E A N L L W R Q E M G G N CTC ATA GAG GCC AAC CTC CTA TGG AGG CAG GAG ATG GGC GGC AAC 1010 T R V E S E N K V V I L D S ATC ACC AGG GTT GAG TCA GAA AAC AAA GTG GTG ATT CTG GAC TCC 1020 F D P L V A E E D E R E I S V TTC GAT CCG CTT GTG GCG GAG GAG GAC GAG CGG GAG ATC TCC GTA 1040 EILRKSRRFAQAL CCC GCA GAA ATC CTG CGG AAG TCT CGG AGA TTC GCC CAG GCC CTG 1050 V W A R P D Y N P P L V E CCC GTT TGG GCG CGG CCG GAC TAT AAC CCC CCG CTA GTG GAG ACG 1070 K K P D Y E P P V V H G C P TGG AAA AAG CCC GAC TAC GAA CCA CCT GTG GTC CAT GGC TGC CCG 1080 K S P P V P P R K K P CTT CCA CCT CCA AAG TCC CCT CCT GTG CCT CCG CCT CGG AAG AAG 1100 T V V L T E S ST L CGG ACG GTG GTC CTC ACT GAA TCA ACC CTA TCT ACT GCC TTG GCC 1110 L A T R S F G S S S T S G GAG CTC GCC ACC AGA AGC TTT GGC AGC TCC TCA ACT TCC GGC ATT 1130 D N T T T S S E P A P S G ACG GGC GAC AAT ACG ACA ACA TCC TCT GAG CCC GCC CCT TCT GGC 1140 S D A E S Y S S M P P D TGC CCC CCC GAC TCC GAC GCT GAG TCC TAT TCC TCC ATG CCC CCC 1160 P G D P D L S D G S W CTG GAG GGG GAG CCT GGG GAT CCG GAT CTT AGC GAC GGG TCA TGG

					E GAG									
					W TGG					V				
					L CTG									
					L TTG					T				
					K									1240 L CTG
					D GAC					v				A GCG
					N		L							1270 S AGC
					S TCA					F				
					H									1300 N AAC
					L CTT					V				
T ACT	T ACC	I ATC	M	132 A GCT	K	N AAC	E GAG	V GTT	F TTC	C TGC	V GTT	Q. CAG	P CCT	1330 E GAG
					P CCA					V				
G	v	R	v	135 C	•	K	М	A	L	Y	D	V	v	1360 T

ĸ	L	P	L	A	v	M	. G	S.	1370 S		G	F	Q	Y
				GCC	GTG								CÃA	
				1380 R	V									1390 S
TCA	CCA	GGA	CAG	CGG	GTT	GAA	ŢTC	CTC			GCG	TGG	AAG	TCC
				M						${f T}$	R	С	F	D
AAG	AAA	ACC	CCA	ATG	GGG	TTC	TCG	TAT	GAT	ACC	CGC	TGC	TTT	GAC
. q	Т	v	T	1410		n	т	R	т	R.	E.	Δ	I	1420 Y
													ATC	-
									1430			•		i.
													K AAG	
				1440)									1450
L		E	R	L	Y	ς π _ι π	G	G	P	L	T	N יייממ	S TCA	
CIC	ACC	GAG	AGG	CII	IAI	GII	GGG	GGC			ACC	M	ICA	AGG
										A			V	
GGG	GAG	AAC	TGC	GGC	TAT	CGC	AĢG	TGC	CGC	GCG	AGC	GGC	GTA	CTG
m	т	C	~	1470 G		m	т.	m		v	т	v	A	1480 R
	_	_											GCC	
	•								149					
													L CTC	
				150			•							1510
С	G	D	D	L	v	V	I	Ċ	Ė	S	A	G	V	Q
TGT	GGC	GAC	GAC	TTA	GTC	GTT	ATC	TGT	GAA	AGC	GCG	GGG	GTC	CAG
F.	מ	A	Δ	s .	T	R	Δ	न	152°		Д	М	Т	R
													ACC	
				153					•					1540
													D GAC	
	-00	-00	-00										-3.3	- -
E	L	I	T	s	С	s	s	N	155 V		V	A	Н	D .
070	CIDC	N m n	202	max.	maa	maa	maa	777	CIDC	mc n	CMC	CCC	CAC	CAC

15/21

1560 1570 G A G K R V Y Y L T R D P T T GGC GCT GGA AAG AGG GTC TAC TAC CTC ACC CGT GAC CCT ACA ACC 1580 PLARAAWETARHTPV... CCC CTC GCG AGA GCT GCG TGG GAG ACA GCA AGA CAC ACT CCA GTC 1590 160 NSWLGNIIMFAPTLW AAT TCC TGG CTA GGC AAC ATA ATC ATG TTT GCC CCC ACA CTG TGG A R M I L M T H F F S V L I A GCG AGG ATG ATA CTG ATG ACC CAT TTC TTT AGC GTC CTT ATA GCC 1620 R D Q L E Q A L D C E I Y G A AGG GAC CAG CTT GAA CAG GCC CTC GAT TGC GAG ATC TAC GGG GCC 1640 C Y S I E P L D L P P I I Q R TGC TAC TCC ATA GAA CCA CTG GAT CTA CCT CCA ATC ATT CAA AGA E I N R V A A C L R K L G V P GAA ATC AAT AGG GTG GCC GCA TGC CTC AGA AAA CTT GGG GTA CCG 1680 1690 P L R A W R H R A R S V R A R CCC TTG CGA GCT TGG AGA CAC CGG GCC CGG AGC GTC CGC GCT AGG L L A R G G R A A I C G K Y L CTT CTG GCC AGA GGC AGG GCT GCC ATA TGT GGC AAG TAC CTC 1710 F N W A V R T K L K L T P I A TTC AAC TGG GCA GTA AGA ACA AAG CTC AAA CTC ACT CCA ATA GCG 1730 A A G Q L D L S G W F T A G Y GCC GCT GGC CAG CTG GAC TTG TCC GGC TGG TTC ACG GCT GGC TAC 1740 S G G D I Y H S V S H A R P R AGC GGG GGA GAC ATT TAT CAC AGC GTG TCT CAT GCC CGG CCC CGC

16/21

1760 W I W F C L L L A A G V G I TGG ATC TGG TTT TGC CTA CTC CTG CTT GCT GCA GGG GTA GGC ATC 1780 ' ' 1770 YLLPNRMSTNPKPQR TAC CTC CTC CCC AAC CGA ATG AGC ACG AAT CCT AAA CCT CAA AGA 1790 K T K R N T N R R P Q D V K F AAG ACC AAA CGT AAC.ACC AAC CGG CGG CCG CAG GAC GTC AAG TTC 1800 P G G G Q I V G G V Y L L P R CCG GGT GGC GGT CAG ATC GTT GGT GGA GTT TAC TTG TTG CCG CGC 1820 R G P R L G V R A T R K AGG GGC CCT AGA TTG GGT GTG CGC GCG ACG AGA AAG ACT TCC GAG 1830 P R G RRQPIPKA CGG TCG CAA CCT CGA GGT AGA CGT CAG CCT ATC CCC AAG GCT CGT 1850 E G R T W A Q P G CGG CCC GAG GGC AGG ACC TGG GCT CAG CCC GGG TAC CCT TGG CCC 1860 L Y G N E G C G W A G W L L S CTC TAT GGC AAT GAG GGC TGC GGG TGG GCG GGA TGG CTC CTG TCT 1880 G S R P S W G P T D P R R CCC CGT GGC TCT CGG CCT AGC TGG GGC CCC ACA GAC CCC CGG CGT 1890 1892 R S R N L G K AGG TCG CGC AAT TTG GGT AAG



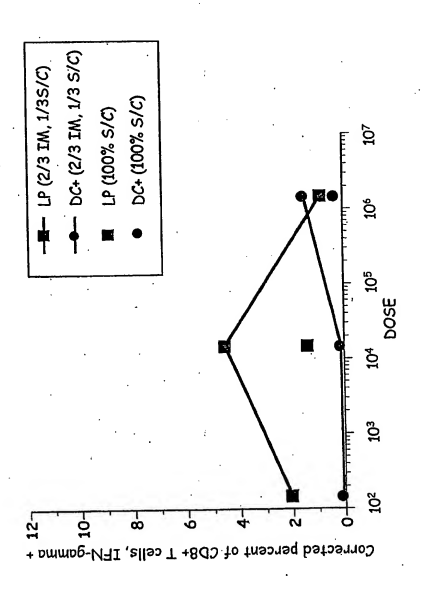


FIG. 6

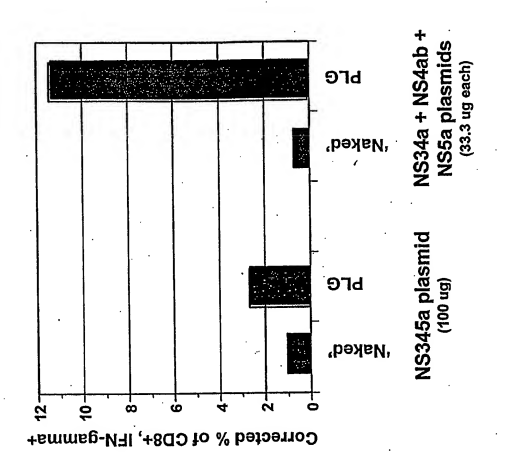
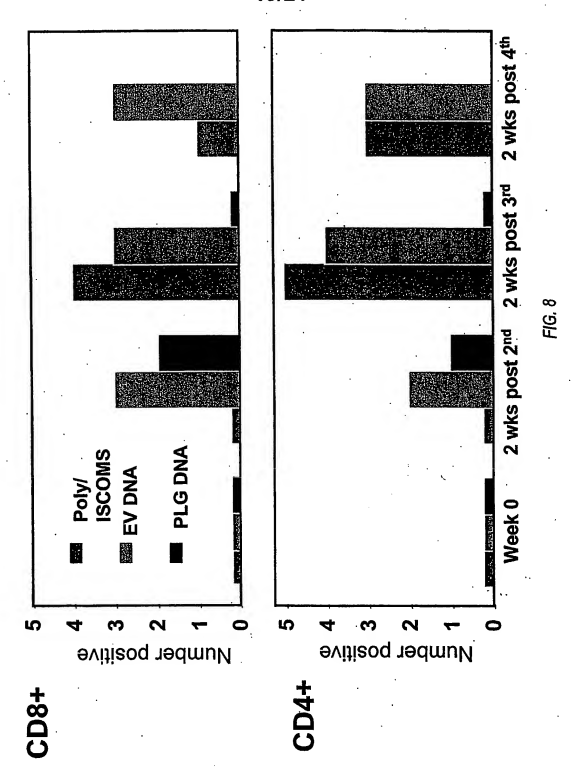
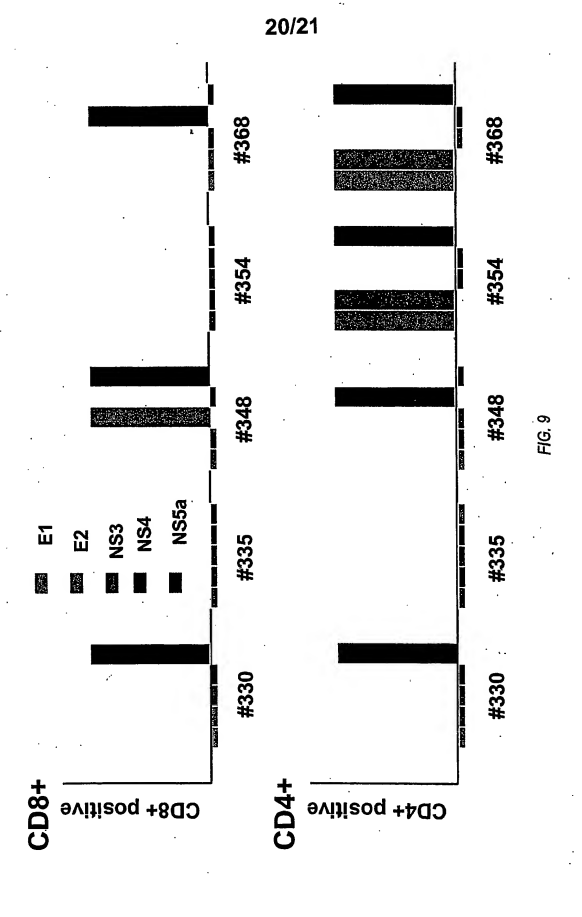


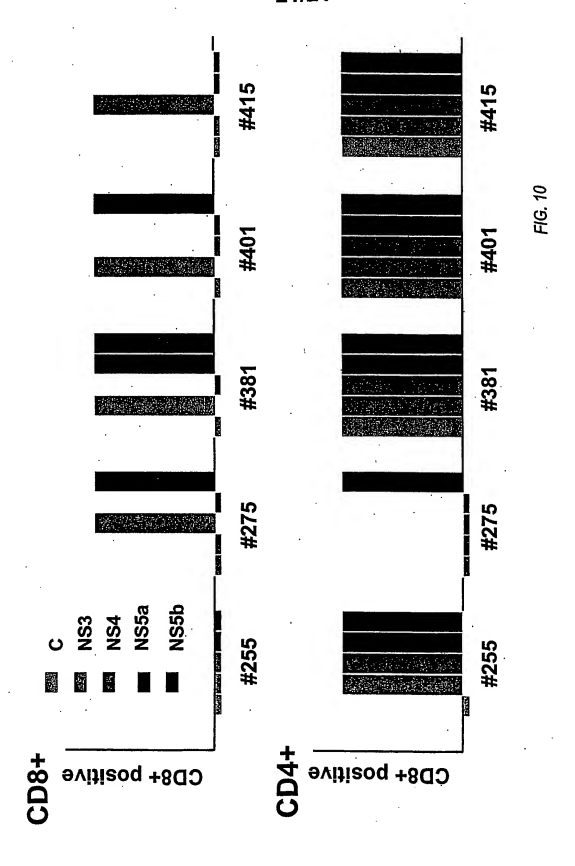
FIG. /











SEQUENCE LISTING

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     SELBY, Mark
     PALIARD, Xavier
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<151> 2002-10-25
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                                                                   48
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aat Asn	gat Asp	tgc Cys 35	cct Pro	aac Asn	tcg Ser	agt Ser	att Ile 40	gtg Val	tac Tyr	gag Glu	gcg Ala	gcc Ala 45	gat Asp	gcc Ala	atc Ile	144
ctg Leu	cac His 50	act Thr	ccg Pro	ggg Gly	tgc Cys	gtc Val 55	cct Pro	tgc Cys	gtt Val	cgc Arg	gag Glu 60	ggc Gly	aac Asn	gcc Ala	tcg Ser	192
agg Arg 65	tgt Cys	tgg Trp	gtg Val	gcg Ala	atg Met 70	acc Thr	cct Pro	acg Thr	gtg Val	gcc Ala 75	acc Thr	agg Arg	gat Asp	ggc	aaa Lys 80	240
ctc Leu	ccc Pro	gcg Ala	acg Thr	cag Gln 85	ctt Leu	cga Arg	cgt Arg	cac His	atc Ile 90	gat Asp	ctg Leu	ctt Leu	gtc Val	ggg Gly 95	agc Ser	288
gcc Ala	acc Thr	ctc Leu	tgt Cys 100	tcg Ser	gcc Ala	ctc Leu	tac Tyr	gtg Val 105	Gly ggg	gac Asp	ctg Leu	tgc Cys	999 Gly 110	tct Ser	gtc Val	336
ttt Phe	ctt Leu	gtc Val 115	ggc	caa Gln	ctg Leu	ttt Phe	acc Thr 120	Phe	tct Ser	ccc Pro	agg Arg	cgc Arg 125	cac His	tgg Trp	acg Thr	384
acg Thr	caa Gln 130	ggt Gly	tgc Cys	aat Asn	tgc Cys	tct Ser 135	atc Ile	tat Tyr	ccc Pro	ggg	cat His 140	Ile	acg Thr	ggt Gly	cac His	432
cgc Arg 145	Met	gca Ala	tgg Trp	gat Asp	atg Met 150	atg Met	atg Met	aac Asn	tgg	Ser 155	Pro	acg Thr	acg Thr	gcg Ala	ttg Leu 160	480
gta Val	atg . Met	gct Ala	cag Gln	ctg Leu 165	Leu	cgg Arg	ato	cca Pro	caa Gln 170	Ala	ato Ile	ttg Leu	gac	atg Met 175	IIe	528
gct Ala	ggt Gly	gct Ala	cac His	Trp	gga Gly	gtc Val	cto Lev	gcg Ala 185	Gly	ata / Ile	a gcg e Ala	tat Tyr	Phe 190	Ser	atg Met	576
gtg Va]	g ggg	aac Asn 195	Trp	gcg Ala	g aag Lys	gtc Val	cto Lev 200	ı Val	gtg . Val	g cto L Lei	g cto 1 Lev	g cta 1 Leu 205	Phe	gcc Ala	ggc Gly	624
gto Va:	gac L Asp 210	Ala	gaa Glu	aco Thi	cac His	gto Val	Thi	ggg Gly	gga Gly	a agi	t gcd r Ala 220	a Gly	cac Hia	c act	gtg Val	672
t c! Se: 22!	r Gly	ttt / Phe	gtt Val	ago Sei	c cto Leu 230	ı Lev	gca 1 Ala	a cca	ggo Gly	gc y Ala 23	а Ьу	g cag s Glr	g aad n Asi	gto n Val	cag Gln 240	720
ct; Le	g ato u Ile	c aac e Asr	e aco	24!	n Gly	agt Ser	tgg Tr	g cad p His	c cto Len 250	u As:	t ago n Se:	c acc	g gc	c cto a Lei 255	g aac 1 Asn 5	768

tgc Cys	aat Asn	gat Asp	agc Ser 260	ctc Leu	aac Asn	acc Thr	ggc Gly	tgg Trp 265	ttg Leu	gca Ala	gly ggg	ctt Leu	ttc Phe 270	tat Tyr	cac His	816
cac His	aag Lys	ttc Phe 275	aac Asn	tct Ser	tca Ser	ggc Gly	tgt Cys 280	cct Pro	gag Glu	agg Arg	cta Leu	gcc Ala 285	agc Ser	tgc Cys	cga Arg	864
ccc Pro	ctt Leu 290	acc Thr	gat Asp	ttt Phe	gac Asp	cagʻ Gln 295	ggc Gly	tgg Trp	ggc Gly	cct Pro	atc Ile 300	agt Ser	tat Tyr	gcc Ala	aac Asn	912
gga Gly 305	agc Ser	ggc Gly	ccc Pro	gac Asp	cag Gln 310	cgc Arg	ccc	tac Tyr	tgc Cys	tgg Trp 315	cac His	tac Tyr	ccc Pro	cca Pro	aaa Lys 320	960
cct Pro	tgc Cys	ggt Gly	att Ile	gtg Val 325	ccc Pro	gcg Ala	aag Lys	Ser	gtg Val 330	tgt Cys	ggt Gly	ccg Pro	gta Val	tat Tyr 335	tgc Cys	1008
ttc Phe	act Thr	ccc Pro	agc Ser 340	ccc Pro	gtg Val	gtg Val	gtg Val	gga Gly 345	acg Thr	acc Thr	gac Asp	agg Arg	tcg Ser 350	ggc Gly	gcg Ala	1056
ccc Pro	acc Thr	tac Tyr 355	agc Ser	tgg Trp	ggt Gly	gaa Glu	aat Asn 360	gat Asp	acg Thr	gac Asp	gtc Val	ttc Phe 365	gtc Val	ctt Leu	aac Asn	1104
aat Asn	acc Thr 370	agg Arg	cca Pro	ccg Pro	ctg Leu	ggc Gly 375	aat Asn	tgg Trp	ttc Phe	ggt Gly	tgt Cys 380	acc Thr	tgg Trp	atg Met	aac Asn	1152
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gjå aaa	gcg Ala	ggc	aac Asn	aac Asn 405	Thr	ctg Leu	cac His	tgc Cys	Pro 410	Thr	gat Asp	tgc Cys	ttc Phe	cgc Arg 415	aag Lys	1248
cat His	ccg Pro	gac Asp	gcc Ala 420		tac Tyr	tct Ser	cgg Arg	tgc Cys 425	Gly	tcc Ser	ggt Gly	ccc Pro	tgg Trp 430	Ile	aca Thr	1296
ccc Pro	agg Arg	tgc Cys 435	Leu	gtc Val	gac Asp	tac Tyr	Pro	Tyr	agg Arg	ctt Leu	tgg Trp	cat His 445	Tyr	cct Pro	tgt Cys	1344
Thr	11e 450	Asn	Туг	Thr	Ile	Phe 455	Lys	Ile	Arg	Met	Tyr 460	Val	Gly	Gly	ytc Val tgc	1392 1440
Gļu 465	His	Arg	Leu	Glu	Ala 470	Ala	Сув	Asn	Trp	475	Arg	Gly	Glu	Arg	480	
Asp	Leu	Glu	Asp	Arg 485	yab	Arg	, Ser	Glu	490	Ser	Pro	Leu	Lev	495		1488
act	aca	cag	tgg	cag	gto	cto	ccg	tgt	tco	tto	aca	acc	cto	, cca	gcc	1536

Thr Thr Gln Trp Gln Val Leu Pro Cys Ser Phe Thr Thr Leu Pro Ala 500 . 505 ttg tcc acc ggc ctc atc cac ctc cac cag aac att gtg gac gtg cag 1584 Leu Ser Thr Gly Leu Ile His Leu His Gln Asn Ile Val Asp Val Gln 520 tac ttg tac ggg gtg ggg tca agc atc gcg tcc tgg gcc att aag tgg Tyr Leu Tyr Gly Val Gly Ser Ser Ile Ala Ser Trp Ala Ile Lys Trp 530 gag tac gtc gtc ctc ctg ttc ctt ctg ctt gca gac gcg cgc gtc tgc 1680 Glu Tyr Val Val Leu Leu Phe Leu Leu Leu Ala Asp Ala Arg Val Cys 555 550 tcc tgc ttg tgg atg atg cta ctc ata tcc caa gcg gaa gcg gct ttg Ser Cys Leu Trp Met Met Leu Leu Ile Ser Gln Ala Glu Ala Ala Leu 570 gag aac ctc gta ata ctt aat gca gca tcc ctg gcc ggg acg cac ggt Glu Asn Leu Val Ile Leu Asn Ala Ala Ser Leu Ala Gly Thr His Gly 580 585 ctt gta tcc ttc ctc gtg ttc ttc tgc ttt gca tgg tat ctg aag ggt 1824 Leu Val Ser Phe Leu Val Phe Phe Cys Phe Ala Trp Tyr Leu Lys Gly 600 aag tgg gtg ccc gga gcg gtc tac acc ttc tac ggg atg tgg cct ctc 1872 Lys Trp Val Pro Gly Ala Val Tyr Thr Phe Tyr Gly Met Trp Pro Leu 615 610 1914 ctc ctg ctc ctg ttg gcg ttg ccc cag cgg gcg tac gcg taa Leu Leu Leu Leu Ala Leu Pro Gln Arg Ala Tyr Ala <210> 4 <211> 637 <212> PRT <213> Artificial Sequence <223> Description of Artificial Sequence: HCV-1 E1/E2/p7 region Ser Phe Ser Ile Phe Leu Leu Ala Leu Leu Ser Cys Leu Thr Val Pro Ala Ser Ala Tyr Gin Val Arg Asn Ser Thr Gly Leu Tyr His Val Thr Asn Asp Cys Pro Asn Ser Ser Ile Val Tyr Glu Ala Ala Asp Ala Ile Leu His Thr Pro Gly Cys Val Pro Cys Val Arg Glu Gly Asn Ala Ser Arg Cys Trp Val Ala Met Thr Pro Thr Val Ala Thr Arg Asp Gly Lys Leu Pro Ala Thr Gln Leu Arg Arg His Ile Asp Leu Leu Val Gly Ser

85 Ala Thr Leu Cys Ser Ala Leu Tyr Val Gly Asp Leu Cys Gly Ser Val 105 Phe Leu Val Gly Gln Leu Phe Thr Phe Ser Pro Arg Arg His Trp Thr 120 Thr Gln Gly Cys Asn Cys Ser Ile Tyr Pro Gly His Ile Thr Gly His Arg Met Ala Trp Asp Met Met Met Asn Trp Ser Pro Thr Thr Ala Leu Val Met Ala Gln Leu Leu Arg Ile Pro Gln Ala Ile Leu Asp Met Ile Ala Gly Ala His Trp Gly Val Leu Ala Gly Ile Ala Tyr Phe Ser Met Val Gly Asn Trp Ala Lys Val Leu Val Val Leu Leu Leu Phe Ala Gly Val Asp Ala Glu Thr His Val Thr Gly Gly Ser Ala Gly His Thr Val Ser Gly Phe Val Ser Leu Leu Ala Pro Gly Ala Lys Gln Asn Val Gln 230 Leu Ile Asn Thr Asn Gly Ser Trp His Leu Asn Ser Thr Ala Leu Asn Cys Asn Asp Ser Leu Asn Thr Gly Trp Leu Ala Gly Leu Phe Tyr His 265 His Lys Phe Asn Ser Ser Gly Cys Pro Glu Arg Leu Ala Ser Cys Arg 280 Pro Leu Thr Asp Phe Asp Gln Gly Trp Gly Pro Ile Ser Tyr Ala Asn 295 Gly Ser Gly Pro Asp Gln Arg Pro Tyr Cys Trp His Tyr Pro Pro Lys 315 Pro Cys Gly Ile Val Pro Ala Lys Ser Val Cys Gly Pro Val Tyr Cys 330 Phe Thr Pro Ser Pro Val Val Val Gly Thr Thr Asp Arg Ser Gly Ala Pro Thr Tyr Ser Trp Gly Glu Asn Asp Thr Asp Val Phe Val Leu Asn Asn Thr Arg Pro Pro Leu Gly Asn Trp Phe Gly Cys Thr Trp Met Asn Ser Thr Gly Phe Thr Lys Val Cys Gly Ala Pro Pro Cys Val Ile Gly

410

Gly Ala Gly Asn Asn Thr Leu His Cys Pro Thr Asp Cys Phe Arg Lys

His Pro Asp Ala Thr Tyr Ser Arg Cys Gly Ser Gly Pro Trp Ile Thr 420 425 430

Pro Arg Cys Leu Val Asp Tyr Pro Tyr Arg Leu Trp His Tyr Pro Cys 445

Thr Ile Asn Tyr Thr Ile Phe Lys Ile Arg Met Tyr Val Gly Gly Val 450 460

Glu His Arg Leu Glu Ala Ala Cys Asn Trp Thr Arg Gly Glu Arg Cys 465 470 475 480

Asp Leu Glu Asp Arg Asp Arg Ser Glu Leu Ser Pro Leu Leu Thr 485 490 490

Thr Thr Gln Trp Gln Val Leu Pro Cys Ser Phe Thr Thr Leu Pro Ala 500 505 510

Leu Ser Thr Gly Leu Ile His Leu His Gln Asn Ile Val Asp Val Gln 515 520 525

Tyr Leu Tyr Gly Val Gly Ser Ser Ile Ala Ser Trp Ala Ile Lys Trp 530 540

Glu Tyr Val Val Leu Leu Phe Leu Leu Leu Ala Asp Ala Arg Val Cys 545 550 560

Ser Cys Leu Trp Met Met Leu Leu Ile Ser Gln Ala Glu Ala Ala Leu 565 570 575

Glu Asn Leu Val Ile Leu Asn Ala Ala Ser Leu Ala Gly Thr His Gly 580 585 590

Leu Val Ser Phe Leu Val Phe Phe Cys Phe Ala Trp Tyr Leu Lys Gly 595 600 605

Lys Trp Val Pro Gly Ala Val Tyr Thr Phe Tyr Gly Met Trp Pro Leu 610 615 620

Leu Leu Leu Leu Ala Leu Pro Gln Arg Ala Tyr Ala 625 630 635

<210> 5

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: consensus sequence

<400> 5

Gly Ser Ala Ala Arg Thr Thr Ser Gly Phe Val Ser Leu Phe Ala Pro 1 5 10 15

Gly Ala Lys Gln Asn 20

<210> 6 <211> 20

<212> DNA <213> Artificial Sequence <223> Description of Artificial Sequence: exemplary CpG oligonucleotide <400> 6 20 tccatgacgt tcctgacgtt <210> 7 <211> 5676 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence: representative NS345Core fusion protein <220> <221> CDS <222> (1)..(5676) atg gct gca tat gca gct cag ggc tat aag gtg cta gta ctc aac ccc Met Ala Ala Tyr Ala Ala Gln Gly Tyr Lys Val Leu Val Leu Asn Pro tct gtt gct gca aca ctg ggc ttt ggt gct tac atg tcc aag gct cat Ser Val Ala Ala Thr Leu Gly Phe Gly Ala Tyr Met Ser Lys Ala His 25 ggg atc gat cct aac atc agg acc ggg gtg aga aca att acc act ggc Gly Ile Asp Pro Asn Ile Arg Thr Gly Val Arg Thr Ile Thr Thr Gly 35 age ecc ate acg tae tee ace tae gge aag tte ett gee gae gge ggg Ser Pro Ile Thr Tyr Ser Thr Tyr Gly Lys Phe Leu Ala Asp Gly Gly 55 tgc tcg ggg ggc gct tat gac ata ata att tgt gac gag tgc cac tcc Cys Ser Gly Gly Ala Tyr Asp Ile Ile Ile Cys Asp Glu Cys His Ser 70 acg gat gcc aca tcc atc ttg ggc att ggc act gtc ctt gac caa gca Thr Asp Ala Thr Ser Ile Leu Gly Ile Gly Thr Val Leu Asp Gln Ala 90 gag act gcg ggg gcg aga ctg gtt gtg ctc gcc acc gcc acc cct ccg 336 Glu Thr Ala Gly Ala Arg Leu Val Val Leu Ala Thr Ala Thr Pro Pro 100 105 ggc tcc gtc act gtg ccc cat ccc aac atc gag gag gtt gct ctg tcc Gly Ser Val Thr Val Pro His Pro Asn Ile Glu Glu Val Ala Leu Ser 115 120 acc acc gga gag atc cct ttt tac ggc aag gct atc ccc ctc gaa gta 432 Thr Thr Gly Glu Ile Pro Phe Tyr Gly Lys Ala Ile Pro Leu Glu Val 135 atc aag ggg ggg aga cat ctc atc ttc tgt cat tca aag aag aag tgc

Ile 145	Lys	Gly	Gly	Arg	His 150	Leu	Ile	Phe	Сув	His 155	Ser	Lys	Lys	ГÀв	Cys 160	
gac Asp	gaa Glu	ctc Leu	gcc Ala	gca Ala 165	aag Lys	ctg Leu	gtc Val	gca Ala	ttg Leu 170	ggc Gly	atc Ile	aat Asn	gcc Ala	gtg Val 175	gcc Ala	528
tac Tyr	tac Tyr	cgc Arg	ggt Gly 180	ctt Leu	gac Asp	gtg Val	tcc Ser	gtc Val 185	atc Ile	ccg Pro	acc Thr	agc Ser	ggc Gly 190	gat Asp	gtt Val	576
gtc Val	gtc Val	gtg Val 195	gca Ala	acc Thr	gat Asp	gcc Ala	ctc Leu 200	atg Met	acc Thr	ggc	tat Tyr	acc Thr 205	ggc	gac Asp	ttc Phe	624
gac Asp	tcg Ser 210	gtg Val	ata Ile	gac Asp	tgc Cys	aat Asn 215	acg Thr	tgt Cys	gtc Val	acc Thr	cag Gln 220	aca Thr	gtc Val	gat Asp	ttc Phe	672
agc Ser 225	ctt Leu	gac Asp	cct Pro	acc Thr	ttc Phe 230	acc Thr	att Ile	gag Glu	aca Thr	atc Ile 235	acg Thr	ctc Leu	ccc Pro	caa Gln	gat Asp 240	720
gct Ala	gtc Val	tcc Ser	cgc Arg	act Thr 245	caa Gln	cgt Arg	cgg Arg	ggc Gly	agg Arg 250	act Thr	ggc	agg Arg	Gly	aag Lys 255	cca Pro	768
ggc	atc Ile	tac Tyr	aga Arg 260	ttt Phe	gtg Val	gca Ala	ccg Pro	999 Gly 265	gag Glu	cgc Arg	ccc Pro	tcc Ser	ggc Gly 270	Met	ttc Phe	816
ga c Asp	tcg Ser	Ser 275	· Val	ctc Leu	tgt Cys	gag Glu	tgc Cys 280	Tyr	gac Asp	gca Ala	ggc	tgt Cys 285	Ala	tgg Trp	tat Tyr	864
gag Glu	cto Leu 290	Thr	r ecc	gcc Ala	gag Glu	act Thr 295	Thr	gtt Val	agg Arg	cta Leu	cga Arg 300	Ala	tac Tyr	atg Met	aac Asn	912
aco Thi	Pro	ggs ggs	g ctt 7 Leu	ccc Pro	gtg Val 310	Cys	cag Gln	gac Asp	cat His	Leu 315	Glu	ttt Phe	tgg Trp	gag Glu	ggc Gly 320	960
gto Val	ttt L Phe	aca Thi	a ggo	cto Leu 325	Thr	cat His	ata Ile	gat Asp	gco Ala 330	His	ttt Phe	cta Leu	tco Ser	cag Gln 335	aca	1008
aag Lys	g cag s Gli	g agt n Sei	ggg Gly 340	/ Glu	aac Asn	ctt Lev	cct Pro	tac Tyr 345	Let	g gta 1 Val	gcg Ala	tac Tyr	caa Glr 350	ı Ala	acc Thr	1056
gt: Va:	g tgo 1 Cy:	gci s Ala 35	a Arg	g gct g Ala	caa Glr	gco Ala	cct Pro 360	Pro	c cca	tco Sei	tgg Tr	gad Asp 369	Glr	g at <u>c</u> n Met	tgg Trp	1104
aa Ly	g tg s Cya 37	s Le	g ati u Il	t cgo	cto J Lev	aag Lys 379	Pro	c acc	c cto	c cat	gg9 Gly 380	Pro	a aca	c Pro	ctg Leu	1152
ct. Le	a ta u Ty	c ag	a cto g Le	g ggd u Gly	gct Ala	gtt a Val	caç L Glı	g aat n Asi	gaa Glu	a ato u Ilo	e acc	c cto	g acg	g cad	c cca Pro	1200

385					390					395					400	
gtc Val	acc Thr	aaa Lys	tac Tyr	atc Ile 405	atg Met	aca Thr	tgc Cys	atg Met	tcg Ser 410	gcc Ala	gac Asp	ctg Leu	gag Glu	gtc Val 415	gtc Val	1248
acg Thr	agc Ser	acc Thr	tgg Trp 420	gtg Val	ctc Leu	gtt Val	ggc	ggc Gly 425	gtc Val	ctg Leu	gct Ala	Ala	ttg Leu 430	gcc Ala	gcg Ala	1296
tat Tyr	tgc Cys	ctg Leu 435	tca Ser	aca Thr	ggc Gly	tgc Cys	gtg Val 440	gtc Val	ata Ile	gtg Val	ggc Gly	agg Arg 445	gtc Val	gtc Val	ttg Leu	1344
tcc Ser	999 Gly 450	aag Lys	ccg Pro	gca Ala	atc Ile	ata Ile 455	cct Pro	gac Asp	agg Arg	gaa Glu	gtc Val 460	ctc Leu	tac Tyr	cga Arg	gag Glu	1392
ttc Phe 465	gat Asp	gag Glu	atg Met	gaa Glu	gag Glu 470	tgc Cys	tct Ser	cag Gln	cac His	tta Leu 475	ccg Pro	tac Tyr	atc Ile	gag Glu	caa Gln 480	1440
61 y 999	atg Met	atg Met	ctc Leu	gcc Ala 485	gag Glu	cag Gln	ttc Phe	aag Lys	cag Gln 490	aag Lys	gcc Ala	ctc Leu	ggc Gly	ctc Leu 495	ctg Leu	1488
cag Gln	acc Thr	gcg Ala	tcc Ser 500	cgt Arg	cag Gln	gca Ala	gag Glu	gtt Val 505	atc Ile	gcc Ala	cct Pro	gct Ala	gtc Val 510	cag Gln	acc Thr	1536
aac Asn	tgg Trp	caa Gln 515	Lys	ctc Leu	gag Glu	acc Thr	ttc Phe 520	tgg Trp	gcg Ala	aag Lys	cat His	atg Met 525	tgg Trp	aac Asn	ttc Phe	1584
atc Ile	agt Ser 530	Gly	ata Ile	caa Gln	tac Tyr	ttg Leu 535	gcg Ala	ggc	ttg Leu	tca Ser	acg Thr 540	ctg Leu	cct Pro	ggt Gly	aac Asn	1632
ccc Pro 545	Ala	att	gct Ala	tca Ser	ttg Leu 550	atg Met	gct Ala	ttt Phe	a ca Thr	gct Ala 555	Ala	gtc Val	acc	agc Ser	Pro 560	1680
cta Leu	acc	act Thr	ago Ser	caa Gln 565	Thr	ct c Leu	ctc Leu	ttc Phe	aac Asn 570	Ile	ttg Leu	gly aaa	Gly 999	tgg Trp 575	gtg Val	1728
gct Ala	gcc	cag Glr	cto Lev 580	ı Ala	gcc Ala	ccc Pro	ggt Gly	gcc Ala 585	Ala	act Thr	gcc Ala	ttt Phe	gtg Val	Gly	gct Ala	1776
ggc Gly	tta Lei	gct Ala 595	Gly	gcc Ala	geo Ala	ato	ggc Gly 600	Ser	gtt Val	gga Gly	a ctg / Lev	999 Gly 605	Lys	gto Val	ctc Leu	1824
ata Ile	gad Asp 610	ıle	ctt E Lei	gca Ala	a ggg a Gly	tat Tyr 615	Gly	Aļā	g ggo	gtg Val	g gcg L Ala 620	Gly	gct Ala	ctt Lev	gtg Val	1872
gca Ala 625	Phe	c aag e Ly:	g ato	e Atg	ago Ser 630	Gly	gag Glu	g gto 1 Val	c cco	s too Ser 63!	r Thi	g gag Glu	ı Ası	cto Lev	g gtc 1 Val 640	1920

aat Asn	cta Leu	ctg Leu	ccc Pro	gcc Ala 645	atc Ile	ctc Leu	tcg Ser	ccc Pro	gga Gly 650	gcc Ala	ctc Leu	gta Val	gtc Val	ggc Gly 655	gtg Val	1968
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gtg Val	cag Gln	tgg Trp 675	atg Met	aac Asn	cgg Arg	ctg Leu	ata Ile 680	gcc Ala	ttc Phe	gcc Ala	tcc Ser	cgg Arg 685	ggg ggg	aac Asn	cat His	2064
gtt Val	tcc Ser 690	ccc Pro	acg Thr	cac His	tac Tyr	gtg Val 695	ccg Pro	gag Glu	agc Ser	gat Asp	gca Ala 700	gct Ala	gcc Ala	cgc Arg	gtc Val	2112
act Thr 705	gcc Ala	ata Ile	ctc Leu	agc Ser	agc Ser 710	ctc Leu	act Thr	gta Val	acc Thr	cag Gln 715	ctc Leu	ctg Leu	agg Arg	cga Arg	ctg Leu 720	2160
cac His	cag Gln	tgg Trp	ata Ile	agc Ser 725	tcg Ser	gag Glu	tgt Cys	acc Thr	act Thr 730	cca Pro	tgc Cys	tcc Ser	ggt Gly	tcc Ser 735	tgg Trp	2208
cta Leu	agg Arg	gac Asp	atc Ile 740	tgg Trp	gac Asp	tgg Trp	ata Ile	tgc Cys 745	gag Glu	gtg Val	ttg Leu	agc Ser	gac Asp 750	ttt Phe	aag Lys	2256
acc Thr	tgg Trp	cta Leu 755	Lys	gct Ala	aag Lys	ctc Leu	atg Met 760	cca Pro	cag Gln	ctg Leu	cct Pro	999 Gly 765	atc Ile	ccc Pro	ttt Phe	2304
gtg Val	tcc Ser 770	Сув	cag Gln	cgc	Gly 999	tat Tyr 775	Lys	G1 ggg	gtc Val	tgg Trp	cga Arg 780	Gly	gac Asp	ggc	atc	2352
atg Met 785	His	act Thr	cgc Arg	tgc Cys	cac His 790	Cys	gga Gly	gćt Ala	gag Glu	atc Ile 795	Thr	gga Gly	cat His	gto Val	aaa Lys 800	2400
aac Asn	ggg Gly	acg Thr	atg Met	agg Arg 805	Ile	gtc Val	ggt	cct Pro	agg Arg 810	Thr	tgo Cys	agg Arg	aac Asn	atg Met 815	tgg Trp	2448
agt Ser	ggg	acc Thr	tto Phe 820	Pro	att Ile	aat Asn	gcc Ala	tac Tyr 825	Thr	acc Thr	ggc Gly	ccc Pro	tgt Cys 830	Thr	ccc Pro	2496
ctt Lev	cct Pro	gcg Ala 835	Pro	g aac Asr	tac Tyr	acc Thr	Phe 840	Ala	j cta Lev	tgg Trp	agg Arg	gte Val 845	Ser	gca Ala	gag Glu	2544
gaa Glu	tac Tyr 850	· Val	g gag L Glu	g ata 1 Ile	agg Arg	Glr 855	ı Val	g ggg	gaq As <u>r</u>	tto Phe	cac His	Tyr	gtg Val	g acc	g ggt Gly	2592
atg Met 865	Thi	act Thi	gac Asp	e aat o Asr	ctt Leu 870	ı Lys	tgo Cys	c ccg	tgo Cyr	cag Glr 879	n Val	c cca	tco Sei	g cco	gaa Glu 880	2640

ttt tto Phe Phe	aca Thr	gaa Glu	ttg Leu 885	gac Asp	G1Y 999	gtg Val	cgc Arg	cta Leu 890	cat His	agg Arg	ttt Phe	gcg Ala	ccc Pro 895	ccc Pro	2688
tgc aag Cys Lys	g ccc s Pro	ttg Leu 900	ctg Leu	cgg Arg	gag Glu	gag Glu	gta Val 905	tca Ser	ttc Phe	aga Arg	gta Val	gga Gly 910	ct c Leu	cac His	2736
gaa ta Glu Ty	c ccg r Pro 915	Val	Gly ggg	tcg Ser	caa Gln	tta Leu 920	cct Pro	tgc Cys	gag Glu	ccc Pro	gaa Glu 925	ccg Pro	gac Asp	gtg Val	2784
gcc gte Ala Va 93	l Leu	acg Thr	tcc Ser	atg Met	ctc Leu 935	act Thr	gat Asp	ccc Pro	tcc Ser	cat His 940	ata Ile	aca Thr	gca Ala	gag Glu	2832
gcg gc Ala Al 945	c ggg a Gly	cga Arg	agg Arg	ttg Leu 950	gcg Ala	agg Arg	gga Gly	tca Ser	ccc Pro 955	ccc Pro	tct Ser	gtg Val	gcc Ala	agc Ser 960	2880
tcc tc Ser Se	g gct r Ala	agc Ser	cag Gln 965	cta Leu	tcc Ser	gct Ala	cca Pro	tct Ser 970	ctc Leu	aag Lys	gca Ala	act Thr	tgc Cys 975	acc Thr	2928
gct aa Ala As	c cat n His	gac Asp 980	Ser	cct Pro	gat Asp	gct Ala	gag Glu 985	ctc Leu	ata Ile	gag Glu	gcc Ala	aac Asn 990	Leu	cta Leu	2976
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aaa gt Lys Va 101	l Val	g att L Ile	ctg Leu	gac Asp	ser 1015	Phe	gat Asp	ccg	Leu	gtg Val	Ala	gag Glu	gag Glu	gac Asp	3072
gag co Glu Ai 1025	gg gag gg Gli	g ato 1 Ile	tcc Ser	gta Val	Pro	gca Ala	gaa Glu	ato Ile	ctg Leu 1035	Arg	j aag j Lys	tct Ser	cgg Arg	aga Arg 1040	3120
ttc go Phe A	cc cag la Gli	g gco n Ala	ctg Leu 1045	Pro	gtt Val	tgg Trp	gcg Ala	Arg 1050	J Pro	gac Asp	tat Tyr	aac Asn	Pro 1055	Pro	3168
cta gi Leu Va	g ga	g acg u Thi 1060	Tr	g aaa b Lys	aag Lys	g ccc s Pro	gac Asp 1065	Ty:	gaa Glu	a cca	a cct	gtg Val	. Val	cat L His	3216
ggc to	gc cc ys Pr 107	o Le	cca Pro	cct Pro	Pro	a aag b Lys 1080	s Sei	c cct	cct Pro	gtg Val	g cct l Pro 1089	Pro	g cct	cgg Arg	3264
aag a Lys L 10	ys Ar	g acg g Thi	g gtg r Val	g gto L Val	c cto L Lev 1099	ı Thi	gaa Glu	tca 1 Sei	a aco	c cta r Len 110	ı Sei	act Thi	gco Ala	ttg a Leu	3312
gcc g Ala G 1105	ag ct lu Le	c gc u Ala	c aco a Thi	c aga r Arg 1110	g Se:	c tti r Phe	t ggo e Gly	age Se:	c tco r Sen 111!	r Se	a act	t too r Sen	c gg c Gl	c att y Ile 1120	3360
acg g	gc ga	c aa	t ac	gac	a ac	a tc	c tc	ga ₉	g cc	c gc	c cc	t tc	t gg	c tgc	3408

-	Asp 2		Thr 7	Thr '	Thr	Ser		Glu 130	Pro	Ala	Pro	Ser (Gly 135	Сув	
ccc ccc Pro Pro	Asp	tcc 9 Ser 1	gac (gct (Ala (gag Glu	Ser	tat Tyr 145	tcc Ser	tcc Ser	atg Met	Pro	ccc Pro 150	ctg Leu	gag Glu	3456
ggg gag Gly Glu	cct Pro 1155	ggg (gat Asp	ccg Pro	Asp	ctt Leu 160	agc Ser	gac Asp	G] À aaa	Ser	tgg Trp .165	tca Ser	acg Thr	gtc Val	3504
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ctg ccc Leu Pro	atc Ile	Asn	gca Ala 205	cta Leu	agc Ser	aac Asn	Ser	ttg Leu 1210	cta Leu	cgt Arg	cac His	His	aat Asn 215	ttg Leu	3648
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aca ttt Thr Phe	gac Asp 1235	aga Arg	ctg Leu	caa Gln	Val	ctg Leu 1240	gac Asp	agc Ser	cat His	Tyr	cag Gln 1245	gac Asp	gta Val	ctc Leu	3744
aag gag Lys Gli 1250	Val د	aaa Lys	gca Ala	Ala	gcg Ala 1255	tca Ser	aaa Lys	gtg Val	ГÄв	gct Ala 1260	Asn	ttg Leu	cta Leu	tcc Ser	3792
gta gag Val Gl 1265	g gaa u Glu	gct. Ala	tgc Cvs	agc Ser	ctg Leu	acg	CCC	cca	cac	tca	gcc	aaa Lvs	tcc Ser	aag	3840
				1270		inr	Pro	Pro	1275	301	1114	-4	•	ьув 1280	
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Phe Gl	y Tyr c atc s Ile	ggg Gly aac Asn 1300	gca Ala 1285 tcc Ser	aaa Lys gtg Val	gac Asp tgg Trp	gtc Val aaa Lys	gac Arg , , gac ; Asp 1305	tgc Cys 1290 ctt Leu	cat His ctg	gcc Ala gaa Glu	aga Arg gac Asp	aag Lys aat Asn 1310	gcc Ala 1295 gta Val	gta Val aca Thr	
phe Gl acc ca Thr Hi	y Tyr c atc s Ile a gac e Asp 1315 g aag u Lys	ggg Gly aac Asn 1300 act	gca Ala 1285 tcc Ser acc Thr	aaa Lys gtg Val atc Ile	gac Asp tgg Trp	gtc Val aaa Lys Ala 1320	gac Asp 1305 aag Lys	tgc Cys 1290 ctt Leu aac Asr	cat His Ctg	gcc Ala gaa Glu gtt Val	aga Arg Asp tto Phe 1325	aag Lys aat Asn 1310 tgc	gcc Ala 1295 gta Val gtt	gta yal aca Thr	3936
Phe Gl acc ca Thr Hi cca at Pro Il cct ga Pro Gl	y Tyr c atc s Ile a gac e Asp 1315 g aag u Lys 0	ggg Gly aac Asn 1300 act Thr	gca Ala 1285 tcc Ser acc Thr	aaa Lys gtg Val atc Ile	gac Asp	gtc Val aaa Lys Ala 1320 g cca g Pro	gac Asp 1305 aag Lys	tgc Cys 1290 ctt Lev d Asr	ctg ctg ctg ctg ctg	gcc Ala gca Ala ggaa gtt ggtt late late late late gtt late late late late late late late la	aga Arg gac Asp ttc Phe 1325	aag Lys aat Asn 1310 c tgc	gcc Ala 1295 gta Val gtt Val	gta yal aca Thr cag Gln gat Asp	3936 3984

1365 1370 1375

	1365	1370	•	
cca gga cag cgg Pro Gly Gln Arg 1380	y Val Glu Phe	ctc gtg caa Leu Val Gln 1385	gcg tgg aag tcc Ala Trp Lys Ser 1390	aag aaa 4176 Lys Lys
acc cca atg ggg Thr Pro Met Gly 1395	Phe Ser Tyr	gat acc cgc Asp Thr Arg 400	tgc ttt gac tcc Cys Phe Asp Ser 1405	aca gtc 4224 Thr Val
act gag agc gad Thr Glu Ser Asp 1410	c atc cgt acg o Ile Arg Thr 1415	gag gag gca Glu Glu Ala	atc tac caa tgt Ile Tyr Gln Cys 1420	tgt gac 4272 Cys Asp
ctc gac ccc cas Leu Asp Pro Gl: 1425	a gcc cgc gtg n Ala Arg Val 1430	Ala Ile Lys	tcc ctc acc gag :Ser Leu Thr Glu 1435	agg ctt 4320 Arg Leu 1440
tat gtt ggg gg Tyr Val Gly Gl	c cct ctt acc y Pro Leu Thr 1445	aat tca agg Asn Ser Arg 1450	ggg gag aắc tgo Gly Glu Asn Cys	ggc tat 4368 Gly Tyr 1455
cgc agg tgc cg Arg Arg Cys Ar 146	g Ala Ser Gly	gta ctg aca Val Leu Thr 1465	act agc tgt ggt Thr Ser Cys Gly 1470	Asn Thr
ctc act tgc ta Leu Thr Cys Ty 1475	r Ile Lys Ala	cgg gca gcc Arg Ala Ala 1480	tgt cga gcc gca Cys Arg Ala Ala 1485	n ggg ctc 4464 n Gly Leu
cag gac tgc ac Gln Asp Cys Th 1490	c atg ctc gtg ar Met Leu Val 1495	Cys Gly Asp	gac tta gtc gtt Asp Leu Val Val 1500	atc tgt 4512 L Ile Cys
gaa agc gcg gg Glu Ser Ala Gl 1505	gg gtc cag gag ly Val Gln Glu 1510	gac gcg gcg Asp Ala Ala	g agc ctg aga gc a Ser Leu Arg Ala 1515	ttc acg 4560 a Phe Thr 1520
gag gct atg ad Glu Ala Met Tl	ec agg tac tec or Arg Tyr Ser 1525	gee eee eet Ala Pro Pro 1530	ggg gac ccc cc Gly Asp Pro Pro O	a caa cca 4608 o Gln Pro 1535
gaa tac gac t Glu Tyr Asp L 15	eu Glu Leu Ile	aca tca tgo Thr Ser Cyr 1545	c tcc tcc aac gt s Ser Ser Asn Va 155	1 Ser val
gcc cac gac g Ala His Asp G 1555	gc gct gga aag ly Ala Gly Lys	agg gtc tag Arg Val Ty: 1560	c tac ctc acc cg r Tyr Leu Thr Ar 1565	t gac cct 4704 g Asp Pro
aca acc ccc c Thr Thr Pro L 1570	tc gcg aga gct eu Ala Arg Ala 1579	a Ala Trp Gl	g aca gca aga ca u Thr Ala Arg Hi 1580	c act cca 4752 s Thr Pro
gtc aat tcc t Val Asn Ser T 1585	gg cta ggc aad rp Leu Gly Asi 1590	c ata atc at n Ile Ile Me	g ttt gcc ccc ac t Phe Ala Pro Th 1595	a ctg tgg 4800 r Leu Trp 1600
gcg agg atg a Ala Arg Met I	ta ctg atg ac le Leu Met Th 1605	c cat ttc tt r His Phe Ph . 161	t agc gtc ctt at e [:] Ser Val Leu Il 0	a gcc agg 4848 e Ala Arg 1615

gac cag ctt gaa Asp Gln Leu Glu 1620	cag gcc ctc q Gln Ala Leu <i>i</i>	gat tgc gag Asp Cys Glu 1625	atc tac ggg gg Ile Tyr Gly A	la Cys Tyr	4896
tcc ata gaa cca Ser Ile Glu Pro 1635	Leu Asp Leu	cct cca atc Pro Pro Ile 640	att caa aga c Ile Gln Arg L 1645	tc cat ggc eu His Gly	4944
ctc agc gca ttt Leu Ser Ala Phe 1650	tca ctc cac Ser Leu His 1655	agt tac tct Ser Tyr Ser	cca ggt gaa a Pro Gly Glu I 1660	tc aat agg le Asn Arg	4992
gtg gcc gca tgc Val Ala Ala Cys 1665	ctc aga aaa Leu Arg Lys 1670	ctt ggg gta Leu Gly Val	ccg ccc ttg c Pro Pro Leu A 1675	ga gct tgg rg Ala Trp 1680	5040
aga cac cgg gcc Arg His Arg Ala	cgg agc gtc Arg Ser Val 1685	cgc gct agg Arg Ala Arg 1690	Leu Leu Ala A	ga gga ggc rg Gly Gly 1695	5088
agg gct gcc ata Arg Ala Ala Ile 1700	tgt ggc aag Cys Gly Lys	tac ctc ttc Tyr Leu Phe 1705	Asn Trp Ala	ta aga aca Val Arg Thr V10	5136
aag ctc aaa ctc Lys Leu Lys Leu 1715	Thr Pro Ile	geg gee get Ala Ala Ala 1720	ggc cag ctg o Gly Gln Leu <i>l</i> 1725	ac ttg tcc Asp Leu Ser	5184
ggc tgg ttc acg Gly Trp Phe Thr 1730	gct ggc tac Ala Gly Tyr 1735	agc ggg ggg Ser Gly Gly	a gac att tat o y Asp Ile Tyr 1 1740	cac agc gtg His Ser Val	5232
tct cat gcc cgg Ser His Ala Arg 1745	ccc cgc tgg Pro Arg Trp 1750	atc tgg tt Ile Trp Ph	t tgc cta ctc (e Cys Leu Leu) 1755	ctg ctt gct Leu Leu Ala 1760	5280
gca ggg gta ggc Ala Gly Val Gly	atc tac ctc Ile Tyr Leu 1765	ctc ccc aa Leu Pro As 177	n Arg Met Ser	acg aat cct Thr Asn Pro 1775	5328
aaa cct caa aga Lys Pro Gln Arg 1780	, Lys Thr Lys	cgt aac ac Arg Asn Th 1785	r Asn Arg Arg	ccg cag gac Pro Gln Asp 790	5376
gtc aag ttc ccg Val Lys Phe Pro 1795	g ggt ggc ggt o Gly Gly Gly	cag atc gt Gln Ile Va 1800	t ggt gga gtt 1 Gly Gly Val 1805	tac ttg ttg Tyr Leu Leu	5424
ccg cgc agg ggo Pro Arg Arg Gly 1810	c cct aga ttg y Pro Arg Leu 1815	Gly Val Ar	gc gcg acg aga gg Ala Thr Arg 1820	aag act tcc Lys Thr Ser	5472
gag cgg tcg caa Glu Arg Ser Gli 1825	a cct cga ggt n Pro Arg Gly 1830	aga cgt ca Arg Arg Gl	ng cct atc ccc In Pro Ile Pro 1835	aag gct cgt Lys Ala Arg 1840	5520
cgg ccc gag gg Arg Pro Glu Gl	c agg acc tgg y Arg Thr Trp 1845	g get cag ed Ala Gln Pr 189	co GIY TYT PTO	tgg ccc ctc Trp Pro Leu 1855	5568

tat ggc aat gag ggc tgc ggg tgg gcg gga tgg ctc ctg tct ccc cgt Tyr Gly Asn Glu Gly Cys Gly Trp Ala Gly Trp Leu Leu Ser Pro Arg 1865 gge tet egg eet age tgg gge eec aca gac eec egg egt agg teg ege Gly Ser Arg Pro Ser Trp Gly Pro Thr Asp Pro Arg Arg Arg Ser Arg 5676 aat ttg ggt aag Asn Leu Gly Lys 1890 <210> 8 <211> 1892 <212> PRT <213> Artificial Sequence <220> <223> Description of Artificial Sequence: representative NS345Core fusion protein Met Ala Ala Tyr Ala Ala Gln Gly Tyr Lys Val Leu Val Leu Asn Pro Ser Val Ala Ala Thr Leu Gly Phe Gly Ala Tyr Met Ser Lys Ala His Gly Ile Asp Pro Asn Ile Arg Thr Gly Val Arg Thr Ile Thr Thr Gly Ser Pro Ile Thr Tyr Ser Thr Tyr Gly Lys Phe Leu Ala Asp Gly Gly 55 Cys Ser Gly Gly Ala Tyr Asp Ile Ile Ile Cys Asp Glu Cys His Ser Thr Asp Ala Thr Ser Ile Leu Gly Ile Gly Thr Val Leu Asp Gln Ala Glu Thr Ala Gly Ala Arg Leu Val Val Leu Ala Thr Ala Thr Pro Pro Gly Ser Val Thr Val Pro His Pro Asn Ile Glu Glu Val Ala Leu Ser 120 Thr Thr Gly Glu Ile Pro Phe Tyr Gly Lys Ala Ile Pro Leu Glu Val Ile Lys Gly Gly Arg His Leu Ile Phe Cys His Ser Lys Lys Lys Cys 155 Asp Glu Leu Ala Ala Lys Leu Val Ala Leu Gly Ile Asn Ala Val Ala Tyr Tyr Arg Gly Leu Asp Val Ser Val Ile Pro Thr Ser Gly Asp Val 185 Val Val Val Ala Thr Asp Ala Leu Met Thr Gly Tyr Thr Gly Asp Phe

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Asp Ser Val Ile Asp Cys Asn Thr Cys Val Thr Gln Thr Val Asp Phe Ser Leu Asp Pro Thr Phe Thr Ile Glu Thr Ile Thr Leu Pro Gln Asp 235 Ala Val Ser Arg Thr Gln Arg Arg Gly Arg Thr Gly Arg Gly Lys Pro Gly Ile Tyr Arg Phe Val Ala Pro Gly Glu Arg Pro Ser Gly Met Phe 265 Asp Ser Ser Val Leu Cys Glu Cys Tyr Asp Ala Gly Cys Ala Trp Tyr 280 Glu Leu Thr Pro Ala Glu Thr Thr Val Arg Leu Arg Ala Tyr Met Asn Thr Pro Gly Leu Pro Val Cys Gln Asp His Leu Glu Phe Trp Glu Gly 315 Val Phe Thr Gly Leu Thr His Ile Asp Ala His Phe Leu Ser Gln Thr 330 Lys Gln Ser Gly Glu Asn Leu Pro Tyr Leu Val Ala Tyr Gln Ala Thr Val Cys Ala Arg Ala Gln Ala Pro Pro Pro Ser Trp Asp Gln Met Trp Lys Cys Leu Ile Arg Leu Lys Pro Thr Leu His Gly Pro Thr Pro Leu Leu Tyr Arg Leu Gly Ala Val Gln Asn Glu Ile Thr Leu Thr His Pro 390 Val Thr Lys Tyr Ile Met Thr Cys Met Ser Ala Asp Leu Glu Val Val 410 Thr Ser Thr Trp Val Leu Val Gly Gly Val Leu Ala Ala Leu Ala Ala Tyr Cys Leu Ser Thr Gly Cys Val Val Ile Val Gly Arg Val Val Leu Ser Gly Lys Pro Ala Ile Ile Pro Asp Arg Glu Val Leu Tyr Arg Glu 455 Phe Asp Glu Met Glu Glu Cys Ser Gln His Leu Pro Tyr Ile Glu Gln 470 Gly Met Met Leu Ala Glu Gln Phe Lys Gln Lys Ala Leu Gly Leu Leu , 490 Gln Thr Ala Ser Arg Gln Ala Glu Val Ile Ala Pro Ala Val Gln Thr 505 Asn Trp Gln Lys Leu Glu Thr Phe Trp Ala Lys His Met Trp Asn Phe 520

The Ser Gly Ile Gln Tyr Leu Ala Gly Leu Ser Thr Leu Pro Gly Asn 530

Pro Ala Ile Ala Ser Leu Met Ala Phe Thr Ala Ala Val Thr Ser Pro 545

Leu Thr Thr Ser Gln Thr Leu Leu Phe Asn Ile Leu Gly Gly Trp Val 575

Ala Ala Gln Leu Ala Ala Pro Gly Ala Ala Thr Ala Phe Val Gly Ala 580

Cly Leu Ala Gly Ala Ala Ile Gly Ser Val Gly Leu Gly Lys Val Leu

Gly Leu Ala Gly Ala Ala Ile Gly Ser Val Gly Leu Gly Lys Val Leu 595 600 605

Ile Asp Ile Leu Ala Gly Tyr Gly Ala Gly Val Ala Gly Ala Leu Val 610 615 620

Ala Phe Lys İle Met Ser Gly Glu Val Pro Ser Thr Glu Asp Leu Val 625 630 630 640

Asn Leu Leu Pro Ala Ile Leu Ser Pro Gly Ala Leu Val Val Gly Val 645 650 . 655

Val Cys Ala Ala Ile Leu Arg Arg His Val Gly Pro Gly Glu Gly Ala 660 665 670

Val Gln Trp Met Asn Arg Leu Ile Ala Phe Ala Ser Arg Gly Asn His
675 680 685

Val Ser Pro Thr His Tyr Val Pro Glu Ser Asp Ala Ala Ala Arg Val 690 695 700

Thr Ala Ile Leu Ser Ser Leu Thr Val Thr Gln Leu Leu Arg Arg Leu 705 710 715 720

His Gln Trp Ile Ser Ser Glu Cys Thr Thr Pro Cys Ser Gly Ser Trp
725 730 735

Leu Arg Asp Ile Trp Asp Trp Ile Cys Glu Val Leu Ser Asp Phe Lys 740 745 750

Thr Trp Leu Lys Ala Lys Leu Met Pro Gln Leu Pro Gly Ile Pro Phe

Val Ser Cys Gln Arg Gly Tyr Lys Gly Val Trp Arg Gly Asp Gly Ile 770 775 780

Met His Thr Arg Cys His Cys Gly Ala Glu Ile Thr Gly His Val Lys 785 790 795 800

Asn Gly Thr Met Arg Ile Val Gly Pro Arg Thr Cys Arg Asn Met Trp

Ser Gly Thr Phe Pro Ile Asn Ala Tyr Thr Thr Gly Pro Cys Thr Pro 820 825 830

Leu Pro Ala Pro Asn Tyr Thr Phe Ala Leu Trp Arg Val Ser Ala Glu 835 840 845

Glu Tyr Val Glu Ile Arg Gln Val Gly Asp Phe His Tyr Val Thr Gly

850 855 860

Met Thr Thr Asp Asn Leu Lys Cys Pro Cys Gln Val Pro Ser Pro Glu 865 870 875 880

- Phe Phe Thr Glu Leu Asp Gly Val Arg Leu His Arg Phe Ala Pro Pro 885 890 895
- Cys Lys Pro Leu Leu Arg Glu Glu Val Ser Phe Arg Val Gly Leu His
- Glu Tyr Pro Val Gly Ser Gln Leu Pro Cys Glu Pro Glu Pro Asp Val 915 920 925
- Ala Val Leu Thr Ser Met Leu Thr Asp Pro Ser His Ile Thr Ala Glu 930 935 940
- Ala Ala Gly Arg Arg Leu Ala Arg Gly Ser Pro Pro Ser Val Ala Ser 945 950 955 960
- Ser Ser Ala Ser Gln Leu Ser Ala Pro Ser Leu Lys Ala Thr Cys Thr 965 970 975
- Ala Asn His Asp Ser Pro Asp Ala Glu Leu Ile Glu Ala Asn Leu Leu 980 985 990
- Trp Arg Gln Glu Met Gly Gly Asn Ile Thr Arg Val Glu Ser Glu Asn 995 1000 1005
- Lys Val Val Ile Leu Asp Ser Phe Asp Pro Leu Val Ala Glu Glu Asp 1010 1015 1020
- Glu Arg Glu Ile Ser Val Pro Ala Glu Ile Leu Arg Lys Ser Arg Arg 1025 1030 1035 1040
- Phe Ala Gln Ala Leu Pro Val Trp Ala Arg Pro Asp Tyr Asn Pro Pro 1045 1050 1055
- Leu Val Glu Thr Trp Lys Lys Pro Asp Tyr Glu Pro Pro Val Val His
- Gly Cys Pro Leu Pro Pro Pro Lys Ser Pro Pro Val Pro Pro Pro Arg 1075 1080 1085
- Lys Lys Arg Thr Val Val Leu Thr Glu Ser Thr Leu Ser Thr Ala Leu 1090 1095 1100
- Ala Glu Leu Ala Thr Arg Ser Phe Gly Ser Ser Ser Thr Ser Gly Ile 1105 1110 1115 1120
- Thr Gly Asp Asn Thr Thr Thr Ser Ser Glu Pro Ala Pro Ser Gly Cys 1125 1130 1135
- Pro Pro Asp Ser Asp Ala Glu Ser Tyr Ser Ser Met Pro Pro Leu Glu 1140 1145 1150
- Gly Glu Pro Gly Asp Pro Asp Leu Ser Asp Gly Ser Trp Ser Thr Val
- Ser Ser Glu Ala Asn Ala Glu Asp Val Val Cys Cys Ser Met Ser Tyr 1170 1175 1180

Ser Trp Thr Gly Ala Leu Val Thr Pro Cys Ala Ala Glu Glu Gln Lys 1185 1190 1195 1200

- Leu Pro Ile Asn Ala Leu Ser Asn Ser Leu Leu Arg His His Asn Leu 1205 1210 1215
- Val Tyr Ser Thr Thr Ser Arg Ser Ala Cys Gln Arg Gln Lys Lys Val
- Thr Phe Asp Arg Leu Gln Val Leu Asp Ser His Tyr Gln Asp Val Leu 1235 1240 1245
- Lys Glu Val Lys Ala Ala Ala Ser Lys Val Lys Ala Asn Leu Leu Ser 1250 1255 1260
- Val Glu Glu Ala Cys Ser Leu Thr Pro Pro His Ser Ala Lys Ser Lys 1265 1270 1275 1280
- Phe Gly Tyr Gly Ala Lys Asp Val Arg Cys His Ala Arg Lys Ala Val 1285 1290 1295
- Thr His Ile Asn Ser Val Trp Lys Asp Leu Leu Glu Asp Asn Val Thr 1300 1305 1310
- Pro Ile Asp Thr Thr Ile Met Ala Lys Asn Glu Val Phe Cys Val Gln 1315 1320 1325
- Pro Glu Lys Gly Gly Arg Lys Pro Ala Arg Leu Ile Val Phe Pro Asp 1330 1335 1340
- Leu Gly Val Arg Val Cys Glu Lys Met Ala Leu Tyr Asp Val Val Thr 1345 1350 1355 1360
- Lys Leu Pro Leu Ala Val Met Gly Ser Ser Tyr Gly Phe Gln Tyr Ser
- Pro Gly Gln Arg Val Glu Phe Leu Val Gln Ala Trp Lys Ser Lys Lys 1380 1385 1390
- Thr Pro Met Gly Phe Ser Tyr Asp Thr Arg Cys Phe Asp Ser Thr Val
- Thr Glu Ser Asp Ile Arg Thr Glu Glu Ala Ile Tyr Gln Cys Cys Asp 1410 1415 1420
- Leu Asp Pro Gln Ala Arg Val Ala Ile Lys Ser Leu Thr Glu Arg Leu 1425 1430 1435 1440
- Tyr Val Gly Gly Pro Leu Thr Asn Ser Arg Gly Glu Asn Cys Gly Tyr 1445 1450 1455
- Arg Arg Cys Arg Ala Ser Gly Val Leu Thr Thr Ser Cys Gly Asn Thr
- Leu Thr Cys Tyr Ile Lys Ala Arg Ala Ala Cys Arg Ala Ala Gly Leu 1475 1480 1485
- Gln Asp Cys Thr Met Leu Val Cys Gly Asp Asp Leu Val Val Ile Cys 1490 1495 1500
- Glu Ser Ala Gly Val Gln Glu Asp Ala Ala Ser Leu Arg Ala Phe Thr

1505 1510 1515 1520 Glu Ala Met Thr Arg Tyr Ser Ala Pro Pro Gly Asp Pro Pro Gln Pro 1525 1530 1535

- Glu Tyr Asp Leu Glu Leu Ile Thr Ser Cys Ser Ser Asn Val Ser Val
- Ala His Asp Gly Ala Gly Lys Arg Val Tyr Tyr Leu Thr Arg Asp Pro 1555 1560 1565
- Thr Thr Pro Leu Ala Arg Ala Ala Trp Glu Thr Ala Arg His Thr Pro 1570 1580
- Val Asn Ser Trp Leu Gly Asn Ile Ile Met Phe Ala Pro Thr Leu Trp 1585 1590 1595 1600
- Ala Arg Met Ile Leu Met Thr His Phe Phe Ser Val Leu Ile Ala Arg 1605 1610 1615
- Asp Gln Leu Glu Gln Ala Leu Asp Cys Glu Ile Tyr Gly Ala Cys Tyr 1620 1625 1630
- Ser Ile Glu Pro Leu Asp Leu Pro Pro Ile Ile Gln Arg Leu His Gly
 1635 1640 1645
- Leu Ser Ala Phe Ser Leu His Ser Tyr Ser Pro Gly Glu Ile Asn Arg 1650 1655 1660
- Val Ala Ala Cys Leu Arg Lys Leu Gly Val Pro Pro Leu Arg Ala Trp 1665 1670 1675 1680
- Arg His Arg Ala Arg Ser Val Arg Ala Arg Leu Leu Ala Arg Gly Gly
 1685 1690 1695
- Arg Ala Ala Ile Cys Gly Lys Tyr Leu Phe Asn Trp Ala Val Arg Thr
- Lys Leu Lys Leu Thr Pro Ile Ala Ala Ala Gly Gln Leu Asp Leu Ser 1715 1720 1725
- Gly Trp Phe Thr Ala Gly Tyr Ser Gly Gly Asp Ile Tyr His Ser Val 1730 1735 1740
- Ser His Ala Arg Pro Arg Trp Ile Trp Phe Cys Leu Leu Leu Leu Ala 1745 1750 1755 1760
- Ala Gly Val Gly Ile Tyr Leu Leu Pro Asn Arg Met Ser Thr Asn Pro 1765 1770 1775
- Lys Pro Gln Arg Lys Thr Lys Arg Asn Thr Asn Arg Arg Pro Gln Asp 1780 1785 1790
- Val Lys Phe Pro Gly Gly Gly Gln Ile Val Gly Gly Val Tyr Leu Leu 1795 1800 1805
- Pro Arg Arg Gly Pro Arg Leu Gly Val Arg Ala Thr Arg Lys Thr Ser 1810 1815 1820
- Glu Arg Ser Gln Pro Arg Gly Arg Arg Gln Pro Ile Pro Lys Ala Arg 1825 1830 1835 1840

Arg Pro Glu Gly Arg Thr Trp Ala Gln Pro Gly Tyr Pro Trp Pro Leu

Tyr Gly Asn Glu Gly Cys Gly Trp Ala Gly Trp Leu Leu Ser Pro Arg Gly Ser Arg Pro Ser Trp Gly Pro Thr Asp Pro Arg Arg Arg Ser Arg 1880 1875 Asn Leu Gly Lys 1890 <210> 9 <211> 546 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence: representative native NS3 protease domain <220> <221> CDS <222> (1)..(546) atg gcg ccc atc acg gcg tac gcc cag cag aca agg ggc ctc cta ggg Met Ala Pro Ile Thr Ala Tyr Ala Gln Gln Thr Arg Gly Leu Leu Gly tgc ata atc acc agc cta act ggc cgg gac aaa aac caa gtg gag ggt Cys Ile Ile Thr Ser Leu Thr Gly Arg Asp Lys Asn Gln Val Glu Gly 20 25 gag gtc cag att gtg tca act gct gcc caa acc ttc ctg gca acg tgc 144 Glu Val Gln Ile Val Ser Thr Ala Ala Gln Thr Phe Leu Ala Thr Cys 35 192 atc aat ggg gtg tgc tgg act gtc tac cac ggg gcc gga acg agg acc Ile Asn Gly Val Cys Trp Thr Val Tyr His Gly Ala Gly Thr Arg Thr 50 atc gcg tca ccc aag ggt cct gtc atc cag atg tat acc aat gta gac Ile Ala Ser Pro Lys Gly Pro Val Ile Gln Met Tyr Thr Asn Val Asp caa gac ctt gtg ggc tgg ccc gct ccg caa ggt agc cga tca ttg aca 288 Gln Asp Leu Val Gly Trp Pro Ala Pro Gln Gly Ser Arg Ser Leu Thr 90 ccc tgc act tgc ggc tcc tcg gac ctt tac ctg gtc acg agg cac gcc 336 Pro Cys Thr Cys Gly Ser Ser Asp Leu Tyr Leu Val Thr Arg His Ala 105 100 gat gtc att ccc gtg cgc cgg cgg ggt gat agc agg ggc agc ctg ctg 384 Asp Val Ile Pro Val Arg Arg Gly Asp Ser Arg Gly Ser Leu Leu tog occ ogg occ att too tac ttg aaa ggo too tog ggg ggt cog otg Ser Pro Arg Pro Ile Ser Tyr Leu Lys Gly Ser Ser Gly Gly Pro Leu

130 135 140

ttg tgc ccc gcg ggg cac gcc gtg ggc ata ttt agg gcc gcg gtg tgc 480 Leu Cys Pro Ala Gly His Ala Val Gly Ile Phe Arg Ala Ala Val Cys 145 150 155 160

acc cgt gga gtg gct aag gcg gtg gac ttt atc cct gtg gag aac cta 528 Thr Arg Gly Val Ala Lys Ala Val Asp Phe Ile Pro Val Glu Asn Leu 165 170 175

gag aca acc atg agg tcc 546
Glu Thr Thr Met Arg Ser
180

<210> 10

<211> 182

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: representative native NS3 protease domain

<400> 10

Met Ala Pro Ile Thr Ala Tyr Ala Gln Gln Thr Arg Gly Leu Leu Gly
1 5 10 15

Cys Ile Ile Thr Ser Leu Thr Gly Arg Asp Lys Asn Gln Val Glu Gly
20 25 30

Glu Val Gln Ile Val Ser Thr Ala Ala Gln Thr Phe Leu Ala Thr Cys 35 40 45

Ile Asn Gly Val Cys Trp Thr Val Tyr His Gly Ala Gly Thr Arg Thr 50 55 60

Ile Ala Ser Pro Lys Gly Pro Val Ile Gln Met Tyr Thr Asn Val Asp 65 70 75 80

Gln Asp Leu Val Gly Trp Pro Ala Pro Gln Gly Ser Arg Ser Leu Thr 85 90 95

Pro Cys Thr Cys Gly Ser Ser Asp Leu Tyr Leu Val Thr Arg His Ala 100 105 110

Asp Val Ile Pro Val Arg Arg Gly Asp Ser Arg Gly Ser Leu Leu 115 , 120 125

Ser Pro Arg Pro Ile Ser Tyr Leu Lys Gly Ser Ser Gly Gly Pro Leu 130 135 140

Leu Cys Pro Ala Gly His Ala Val Gly Ile Phe Arg Ala Ala Val Cys 145 150 155 160

Thr Arg Gly Val Ala Lys Ala Val Asp Phe Ile Pro Val Glu Asn Leu 165 170 175

Glu Thr Thr Met Arg Ser 180